

```
* The Help Desk staff at this number will handle all APS
* related questions.
* * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * *
* >>>>>>>>>>>> NEW SUNDAY HOURS !!! <<<<<<<<<<<<
*
* The APS is available:
*           6:30am - 9:00pm Monday through Friday
*           7:30am - 5:00pm Saturday, Sunday, Holidays
*
* APS is unavailable Thanksgiving Day, Christmas Day,
* and New Year's Day.
*
* * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * * *
FILE 'USPAT' ENTERED AT 14:25:39 ON 14 MAY 1998
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FILE 'USPAT' ENTERED AT 14:25:39 ON 14 MAY 1998

=> s cd4(P) (antibod?) (P) (autoimmun? or arthritis or sclerosis)

2027 CD4
28008 ANTIBOD?
4066 AUTOIMMUN?
10097 ARTHRITIS
3197 SCLEROSIS
L1 75 CD4 (P) (ANTIBOD?) (P) (AUTOIMMUN? OR ARTHRITIS OR SCLEROSIS)

=> s 11/clm

192 CD4/CLM
7759 ANTIBOD?/CLM
385 AUTOIMMUN?/CLM
882 ARTHRITIS/CLM
296 SCLEROSIS/CLM
3 (CD4/CLM(P) (ANTIBOD?/CLM) (P) (AUTOIMMUN?/CLM OR ARTHRITIS/CLM
R SCLEROSIS/CLM))

=> d 12 1-3

1. 5,750,105, May 12, 1998, Recombinant antibodies for human therapy; Roland A. Newman, et al., 424/133.1, 137.1, 138.1, 177.1; 530/387.3 [IMAGE AVAILABLE]
2. 5,741,488, Apr. 21, 1998, Treatment of rheumatoid arthritis with anti-CD4 antibodies in conjunction with anti-TNF antibodies; Marc Feldman, et al., 424/154.1, 130.1, 141.1, 143.1, 144.1, 145.1, 153.1, 158.1, 173.1 [IMAGE AVAILABLE]
3. 4,695,459, Sep. 22, 1987, Method of treating autoimmune diseases that are mediated by Leu3/CD4 phenotype T cells; Lawrence Steinman, et al., 424/154.1, 173.1, 810; 514/825, 863, 866, 885; 530/388.75, 868; 935/107 [IMAGE AVAILABLE]

=> d 12 1-3 kwic

• US PAT NO: 5,750,105 [IMAGE AVAILABLE]

L2: 1 of 3

CLAIMS:

CLMS (8)

8. The method of claim 1, wherein the treatment comprises treatment of a disease selected from the group consisting of rheumatoid **arthritis**, eczema, and immuno-modulated diseases, and the antigen bound by the **antibody** is **CD4**.

US PAT NO: 5,741,488 [IMAGE AVAILABLE]

L2: 2 of 3

CLAIMS:

CLMS (1)

We claim:

1. A method of treating rheumatoid **arthritis** in a mammal comprising administering to said mammal a therapeutically effective amount of **anti-CD4 antibody** and **anti-TNF antibody**.

US PAT NO: 4,695,459 [IMAGE AVAILABLE]

L2: 3 of 3

CLAIMS:

CLMS (1)

We claim:

1. A method of treating a patient for an **autoimmune** disease that is mediated by Leu3(**CD4**) phenotype T cells comprising parenterally administering a therapeutically effective amount of an anti-Leu3(**CD4**) **antibody** that binds to said T cells to the patient.

=> d 13 1-3 date

'L3' NOT FOUND

=> d 12 1-3 date

L2: 1 of 3

TITLE: Recombinant antibodies for human therapy

US PAT NO: 5,750,105 DATE ISSUED: May 12, 1998
[IMAGE AVAILABLE]

APPL-NO: 08/476,349 DATE FILED: Jun. 7, 1995

REL-US-DATA: Division of Ser. No. 379,072, Dec. 5, 1995, which is a continuation of Ser. No. 912,292, Jul. 10, 1992, abandoned, which is a continuation-in-part of Ser. No. 856,281, Mar. 23, 1992, abandoned, which is a continuation-in-part of Ser. No. 735,064, Jul. 25, 1991, abandoned.

L2: 2 of 3

TITLE: Treatment of rheumatoid arthritis with anti-CD4 antibodies in conjunction with anti-TNF antibodies

US PAT NO: 5,741,488 DATE ISSUED: Apr. 21, 1998
[IMAGE AVAILABLE]

APPL-NO: 08/403,785 DATE FILED: May 3, 1995

PCT-NO: PCT/GB93/02070 PCT-FILED: Oct. 6, 1993
371-DATE: May 3, 1995

L2: 3 of 3

TITLE: Method of treating autoimmune diseases that are mediated by Leu3/CD4 phenotype T cells

US PAT NO: 4,695,459 DATE ISSUED: Sep. 22, 1987
[IMAGE AVAILABLE]

APPL-NO: 06/686,126 DATE FILED: Dec. 26, 1984

=> d his

(FILE 'USPAT' ENTERED AT 14:25:39 ON 14 MAY 1998)

L1 75 S CD4(P) (ANTIBOD?) (P) (AUTOIMMUN? OR ARTHRITIS OR SCLEROSIS
)

L2 3 S L1/CLM

=> d 11 1-75

1. 5,750,332, May 12, 1998, Peptomers with enhanced immunogenicity; Frank A. Robey, et al., 435/5, 974; 514/2, 13 [IMAGE AVAILABLE]
2. 5,750,105, May 12, 1998, Recombinant antibodies for human therapy; Roland A. Newman, et al., 424/133.1, 137.1, 138.1, 177.1; 530/387.3 [IMAGE AVAILABLE]
3. 5,747,265, May 5, 1998, Method for measuring the amount of a cell-associated molecule; George H. Parsons, et al., 435/7.2, 7.24 [IMAGE AVAILABLE]
4. 5,747,036, May 5, 1998, Methods and compositions for detecting and treating a subset of human patients having an autoimmune disease; Michael Brenner, et al., 424/144.1, 154.1, 173.1, 178.1 [IMAGE AVAILABLE]
5. 5,741,899, Apr. 21, 1998, Chimeric receptors comprising janus kinase for regulating cellular pro lification; Daniel J. Capon, et al., 536/23.4; 435/69.7, 320.1, 325, 377; 530/350, 387.3 [IMAGE AVAILABLE]
6. 5,741,488, Apr. 21, 1998, Treatment of rheumatoid **arthritis** with **anti-CD4 antibodies** in conjunction with **anti-TNF antibodies**; Marc Feldman, et al., 424/154.1, 130.1, 141.1, 143.1, 144.1, 145.1, 153.1, 158.1, 173.1 [IMAGE AVAILABLE]
7. 5,736,138, Apr. 7, 1998, Monoclonal antibodies with specific binding against membrane proteins on human cells, and pharmaceutical compositions containing them; Klaus Pfizenmaier, et al., 424/143.1, 133.1, 144.1, 152.1, 154.1, 172.1, 173.1, 809; 435/70.21; 530/351, 387.1, 388.22, 388.73, 388.85, 388.9, 399, 866 [IMAGE AVAILABLE]
8. 5,734,023, Mar. 31, 1998, MHC class II .beta. chain/peptide complexes useful in ameliorating deleterious immune responses; Bishwajit Nag, et al., 530/403; 424/185.1, 193.1; 530/300, 395, 402, 868 [IMAGE AVAILABLE]
9. 5,728,680, Mar. 17, 1998, Methods for normalizing numbers of lymphocytes; Vyacheslav G. Morozov, et al., 514/19, 9, 11 [IMAGE AVAILABLE]
10. 5,728,533, Mar. 17, 1998, Human .beta..sub.2 integrin .alpha..subunit; W. Michael Gallatin, et al., 435/7.1, 7.8; 530/350, 380 [IMAGE AVAILABLE]
11. 5,723,503, Mar. 3, 1998, Biological treatment for rheumatoid arthritis; J. Bruce Smith, et al., 424/93.1, 93.71, 534 [IMAGE AVAILABLE]

12. 5,718,883, Feb. 17, 1998, Transgenic animal model for autoimmune diseases; David M. Harlan, et al., 424/9.2; 435/172.3; 514/2; 800/2, DIG.1 [IMAGE AVAILABLE]

13. 5,714,350, Feb. 3, 1998, Increasing antibody affinity by altering glycosylation in the immunoglobulin variable region; Man Sung Co, et al., 435/69.6; 424/133.1; 435/70.21, 71.1, 172.1; 530/387.3; 935/49, 50 [IMAGE AVAILABLE]

14. 5,712,149, Jan. 27, 1998, Chimeric receptor molecules for delivery of co-stimulatory signals; Margo R. Roberts, 435/252.3, 69.7, 320.1; 530/350; 536/23.4 [IMAGE AVAILABLE]

15. 5,710,257, Jan. 20, 1998, Method of causing selective immunosuppression using HL-60-related lectins; Jeffrey J. Seilhamer, et al., 530/396; 435/172.3; 530/350 [IMAGE AVAILABLE]

16. 5,707,626, Jan. 13, 1998, Methods of treating HIV infection using antibodies to the U2 small nuclear ribonuclear protein; Angeline Douvas, et al., 424/160.1, 148.1, 152.1, 172.1 [IMAGE AVAILABLE]

17. 5,705,732, Jan. 6, 1998, Universal donor cells; Peter J. Sims, et al., 800/2; 435/172.3; 536/23.1; 800/DIG.1 [IMAGE AVAILABLE]

18. 5,696,237, Dec. 9, 1997, Recombinant antibody-toxin fusion protein; David FitzGerald, et al., 530/387.3, 388.22, 391.7 [IMAGE AVAILABLE]

19. 5,693,780, Dec. 2, 1997, Recombinant antibodies for human therapy; Roland A. Newman, et al., 536/23.53; 435/252.3, 320.1 [IMAGE AVAILABLE]

20. 5,693,760, Dec. 2, 1997, Method of causing selective immunosuppression using HL-60 related lectins; Jeffrey J. Seilhamer, et al., 530/396; 424/278.1; 435/172.3; 530/350, 827 [IMAGE AVAILABLE]

21. 5,693,617, Dec. 2, 1997, Inhibitors of the 26s proteolytic complex and the 20s proteasome contained therein; Ross L. Stein, et al., 514/18, 19; 530/331; 560/20, 27, 31, 32, 41, 47, 159 [IMAGE AVAILABLE]

22. 5,690,933, Nov. 25, 1997, Monoclonal antibodies for inducing tolerance; Stephen Paul Cobbold, et al., 424/144.1, 143.1, 153.1, 154.1, 173.1; 514/11 [IMAGE AVAILABLE]

23. 5,686,281, Nov. 11, 1997, Chimeric receptor molecules for delivery of co-stimulatory signals; Margo R. Roberts, 435/172.3, 7.1, 7.2, 69.7; 536/23.4 [IMAGE AVAILABLE]

24. 5,681,722, Oct. 28, 1997, Recombinant antibodies for human therapy; Roland A. Newman, et al., 435/69.7, 6, 91.2; 530/387.3; 536/23.53, 24.33 [IMAGE AVAILABLE]

25. 5,675,060, Oct. 7, 1997, Transgenic arthritic mice expressing a T-cell receptor transgene; Christophe O. Benoist, et al., 800/2; 424/9.2 [IMAGE AVAILABLE]

26. 5,674,692, Oct. 7, 1997, Methods for diabetes susceptibility assessment in asymptomatic patients; Steinunn Baekkeskov, et al., 435/7.21, 7.4; 436/506, 518 [IMAGE AVAILABLE]

27. 5,674,487, Oct. 7, 1997, Method for treating autoimmune diseases; J. Bruce Smith, et al., 424/93.71, 93.7 [IMAGE AVAILABLE]

28. 5,670,324, Sep. 23, 1997, Use of chimeric CD4-src protein tyrosine kinases in drug screening and detection of an immune response; Dan Littman, et al., 435/6, 15, 69.7 [IMAGE AVAILABLE]

29. 5,670,150, Sep. 23, 1997, Non-depleting CD4-specific monoclonal antibodies for the treatment of insulin-dependent diabetes mellitus (IDDM); Anne Cooke, et al., 424/154.1, 143.1, 145.1, 158.1; 530/388.24, 388.75 [IMAGE AVAILABLE]

30. 5,667,967, Sep. 16, 1997, T-cell receptor variable transcripts as disease related markers; Lawrence Steinman, et al., 435/6, 91.2; 935/77, 78 [IMAGE AVAILABLE]

31. 5,665,772, Sep. 9, 1997, O-alkylated rapamycin derivatives and their use, particularly as immunosuppressants; Sylvain Cottens, et al., 514/514; 540/456 [IMAGE AVAILABLE]

32. 5,665,764, Sep. 9, 1997, Tricyclic inhibitors of matrix metalloproteinases; Donald Hupe, et al., 514/468; 549/460, 461 [IMAGE AVAILABLE]

33. 5,658,745, Aug. 19, 1997, Cell enumeration immunoassay; Richard Alfred Greene, et al., 435/7.24; 424/154.1, 534; 435/7.92, 7.95, 967, 974; 436/63, 172, 524, 531, 541, 546, 548, 811 [IMAGE AVAILABLE]

34. 5,658,570, Aug. 19, 1997, Recombinant antibodies for human therapy; Roland A. Newman, et al., 424/184.1; 435/69.6, 70.21, 172.2, 172.3; 530/388.22; 935/96 [IMAGE AVAILABLE]

35. 5,639,869, Jun. 17, 1997, Mycoplasma arthritidis T-cell mitogen; Barry C. Cole, et al., 536/23.7; 424/264.1; 530/326, 350, 825 [IMAGE AVAILABLE]

36. 5,635,599, Jun. 3, 1997, Fusion proteins comprising circularly permuted ligands; Ira H. Pastan, et al., 530/351; 435/69.1, 69.5, 69.52, 69.7, 172.3; 530/350 [IMAGE AVAILABLE]

37. 5,627,206, May 6, 1997, Tricyclic inhibitor of matrix metalloproteinases; Donald Hupe, et al., 514/468; 549/461 [IMAGE AVAILABLE]

38. 5,627,035, May 6, 1997, Peptides that block human immunodeficiency virus and methods of use thereof; Anders Vahine, et al., 435/7.2; 424/188.1; 530/327, 328, 329, 330 [IMAGE AVAILABLE]

39. 5,626,843, May 6, 1997, Treatment of autoimmune diseases, including AIDS, by removal of interferons, TNFs and receptors therefor; Simon V. Skurkovich, et al., 424/140.1; 604/6 [IMAGE AVAILABLE]

40. 5,624,895, Apr. 29, 1997, Treatment and/or prevention of type I diabetes mellitus with gamma interferon administration; Douglas Sobel, 514/8; 424/85.1, 85.2, 85.4, 85.5, 85.6, 85.7; 514/866 [IMAGE AVAILABLE]

41. 5,622,853, Apr. 22, 1997, T lymphocyte precursor; Leon W. M. M. Terstappen, et al., 435/372.3, 2, 7.2, 325 [IMAGE AVAILABLE]

42. 5,620,889, Apr. 15, 1997, Human anti-Fas IgG1 monoclonal antibodies; David H. Lynch, et al., 435/332; 424/144.1; 435/334, 343.2; 530/387.1, 388.2, 388.23, 388.24, 388.75 [IMAGE AVAILABLE]

43. 5,616,458, Apr. 1, 1997, Tripterygium wilfordii hook F extracts and components, and uses thereof; Peter E. Lipsky, et al., 435/4; 424/78.05, 195.1; 435/7.5, 7.9; 514/469, 821, 825, 886 [IMAGE AVAILABLE]

44. 5,614,192, Mar. 25, 1997, T cell receptor peptides as therapeutics for immune-related disease; Arthur A. Vandenbark, 424/185.1, 184.1, 193.1; 514/2, 12, 16; 530/300, 324, 328, 868 [IMAGE AVAILABLE]

45. 5,602,095, Feb. 11, 1997, Recombinant *pseudomonas* exotoxin with increased activity; Ira H. Pastan, et al., 514/12; 424/192.1, 193.1, 236.1; 435/69.1, 69.3, 69.7, 172.3, 252.3, 252.33, 320.1; 514/2; 530/350, 351, 403, 825; 930/200 [IMAGE AVAILABLE]

46. 5,583,153, Dec. 10, 1996, Use of taxol in the treatment of rheumatoid arthritis; Ernest Brahn, 514/449, 475 [IMAGE AVAILABLE]

47. 5,583,033, Dec. 10, 1996, T lymphocyte precursor; Leon W. M. M. Terstappen, et al., 435/7.21, 7.24, 378 [IMAGE AVAILABLE]

48. 5,580,772, Dec. 3, 1996, Association between a novel human intracisternal A-type retroviral particle-type II (HIAP-II) and idiopathic CD4+ T-lymphocytopenia (ICL); Robert F. Garry, Jr., 435/235.1; 424/207.1; 435/5, 239 [IMAGE AVAILABLE]

49. 5,580,562, Dec. 3, 1996, Preparations and uses thereof for immunosuppression; Peter E. Lipsky, et al., 424/195.1; 514/885, 908; 549/228, 297, 298 [IMAGE AVAILABLE]

50. 5,571,507, Nov. 5, 1996, Methods of treating diabetes; Vicki E. Rubin-Kelley, et al., 424/85.2; 514/866; 530/321, 351 [IMAGE AVAILABLE]

51. 5,556,754, Sep. 17, 1996, Methods for assessing the ability of a candidate drug to suppress MHC class I expression; Dinah S. Singer, et al., 435/6, 91.1; 436/63, 501; 536/24.31, 24.33; 935/34, 36, 77, 78 [IMAGE AVAILABLE]

52. 5,550,132, Aug. 27, 1996, Hydroxyalkylammonium-pyrimidines or purines and nucleoside derivatives, useful as inhibitors of inflammatory cytokines; Bradley J. Benson, et al., 514/269, 274; 544/311, 312, 313, 314 [IMAGE AVAILABLE]

53. 5,545,716, Aug. 13, 1996, Superantigen agonist and antagonist peptides; Howard M. Johnson, et al., 530/324, 325, 326 [IMAGE AVAILABLE]

54. 5,538,854, Jul. 23, 1996, Method for the determination of predisposition to autoimmune disease; Denise Faustman, 435/7.24, 6; 436/86, 506, 516 [IMAGE AVAILABLE]

55. 5,521,288, May 28, 1996, CD28IG fusion protein; Peter S. Linsley, et al., 530/387.3; 435/7.2, 7.92, 69.1, 69.7, 91.1, 252.3, 252.33, 320.1; 530/300, 350, 387.1, 395, 409, 866, 867, 868; 536/23.1, 23.4, 23.53 [IMAGE AVAILABLE]

56. 5,519,114, May 21, 1996, Retroviral superantigens, superantigen peptides, and methods of use; Howard M. Johnson, et al., 530/324; 424/188.1, 278.1; 435/5; 930/221 [IMAGE AVAILABLE]

57. 5,514,661, May 7, 1996, Immunological activity of rhamnolipids; Goran Piljac, et al., 514/25, 814, 861, 863, 864, 878, 883, 885, 886, 887, 889, 903, 908 [IMAGE AVAILABLE]

58. 5,504,000, Apr. 2, 1996, Chimeric protein tyrosine kinases; Dan Littman, et al., 435/194, 69.1, 69.7; 530/350; 536/22.1, 23.1, 23.2, 23.4, 23.5 [IMAGE AVAILABLE]

59. 5,500,340, Mar. 19, 1996, Inhibition of IL-2 production by *Tripterygium wilfordii* Hook F extract; Peter E. Lipsky, et al., 435/6; 436/63; 935/34, 77 [IMAGE AVAILABLE]

60. 5,468,481, Nov. 21, 1995, MHC class II-peptide conjugates useful in ameliorating autoimmunity; Somesh D. Sharma, et al., 424/185.1, 184.1, 193.1, 278.1; 514/2, 8; 530/395, 402, 403, 868 [IMAGE AVAILABLE]

61. 5,466,675, Nov. 14, 1995, Immunological activity of rhamnolipids; Goran Piljac, et al., 514/25, 814, 861, 863, 864, 878, 883, 885, 886, 887, 889, 903, 908 [IMAGE AVAILABLE]

62. 5,445,940, Aug. 29, 1995, Methods and compositions for detecting and treating a subset of human patients having an autoimmune disease; Michael B. Brenner, et al., 435/7.24, 6; 436/501, 506, 512, 548 [IMAGE AVAILABLE]

63. 5,439,819, Aug. 8, 1995, Chimeric protein tyrosine kinases; Dan Littman, et al., 435/372.3, 69.1, 69.7, 194; 530/350; 536/22.1, 23.1, 23.2, 23.4, 23.5 [IMAGE AVAILABLE]

64. 5,397,702, Mar. 14, 1995, Assay for and treatment of autoimmune diseases; Michael D. Cahalan, et al., 435/69.1, 6, 172.3; 536/23.1, 23.5, 25.5 [IMAGE AVAILABLE]

65. 5,294,443, Mar. 15, 1994, Tripterygium wilford II hook f extracts and components thereof for immunosuppression; Peter E. Lipsky, et al., 424/195.1; 514/885 [IMAGE AVAILABLE]

66. 5,284,935, Feb. 8, 1994, MHC-mediated toxic conjugates useful in ameliorating autoimmunity; Brian R. Clark, et al., 424/185.1, 193.1, 810; 530/395, 403, 806, 807, 868 [IMAGE AVAILABLE]

67. 5,270,199, Dec. 14, 1993, Human mannose-binding protein; Raymond A. B. Ezekowitz, 435/372.1, 69.1, 172.3, 235.1, 252.3, 252.33, 254.11, 254.2, 320.1; 530/350; 536/23.4, 23.5; 935/18, 27, 32, 34, 38, 55, 62, 70, 72 [IMAGE AVAILABLE]

68. 5,260,422, Nov. 9, 1993, MHC conjugates useful in ameliorating autoimmunity; Brian R. Clark, et al., 424/185.1, 193.1, 810; 530/402, 403, 868 [IMAGE AVAILABLE]

69. 5,252,556, Oct. 12, 1993, Fragment capable of binding anti-CD43 autoantibodies; Blair Ardmann, 424/185.1; 435/69.1, 69.3; 514/8; 530/350, 395 [IMAGE AVAILABLE]

70. 5,246,970, Sep. 21, 1993, Method of inhibiting nitric oxide formation; Joseph R. Williamson, et al., 514/632, 903 [IMAGE AVAILABLE]

71. 5,223,426, Jun. 29, 1993, Monoclonal antibodies reactive with defined regions of the T-cell antigen receptor; Robert V. Skibbens, et al., 435/331; 424/144.1, 154.1; 530/387.1, 387.9, 388.22, 388.75 [IMAGE AVAILABLE]

72. 5,194,425, Mar. 16, 1993, MHC-mediated toxic conjugates useful in ameliorating autoimmunity; Somesh D. Sharma, et al., 424/193.1, 185.1; 514/8, 903; 530/395, 402, 403 [IMAGE AVAILABLE]

73. 5,158,884, Oct. 27, 1992, Immunodominant acetylcholine receptor peptides useful for T-helper cell sensitization; Bianca M. Conti-Tronconi, et al., 435/331; 530/326 [IMAGE AVAILABLE]

74. 5,130,297, Jul. 14, 1992, Conjugates useful in ameliorating autoimmunity MHC-II-peptide; Somesh D. Sharma, et al., 514/8, 825, 903; 530/395, 403, 838 [IMAGE AVAILABLE]

75. 4,695,459, Sep. 22, 1987, Method of treating autoimmune diseases that are mediated by Leu3/CD4 phenotype T cells; Lawrence Steinman, et al., 424/154.1, 173.1, 810; 514/825, 863, 866, 885; 530/388.75, 868; 935/107 [IMAGE AVAILABLE]

ber

DIALOG INFORMATION SERVICES

PLEASE LOGON:

ENTER PASSWORD:

Op58093fe

Welcome to DIALOG

Dialog level 98.04.30D

Last logoff: 18may98 14:04:41

Logon file001 18may98 17:20:42

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*** File 728 is not working. ***

File 1:ERIC 1966-1998/Mar

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Set	Items	Description
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? b 410

18may98 17:20:47	User208760	Session D1032.1
\$0.03	0.001	Hrs File1
\$0.03	Estimated cost File1	
\$0.03	Estimated cost this search	
\$0.03	Estimated total session cost 0.001 Hrs.	

File 410:Chronolog(R) 1981-1998/May

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Set	Items	Description
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? set hi ;set hi

HIGHLIGHT set on as ''

HIGHLIGHT set on as ''

? begin 55,72,154,399,351

18may98 17:21:02	User208760	Session D1032.2
\$0.00	0.004	Hrs File410
\$0.00	Estimated cost File410	
\$0.01	FTSNET	
\$0.01	Estimated cost this search	
\$0.04	Estimated total session cost 0.005 Hrs.	

SYSTEM:OS - DIALOG OneSearch

File 55:BIOSIS PREVIEWS(R) 1985-1998/May W2

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File 72:EMBASE 1985-1998/May W2

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File 154:MEDLINE(R) 1985-1998/Jul W2

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File 399:CA SEARCH(R) 1967-1998/UD=12820
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*File 399: Use is subject to the terms of your user/customer agreement.
RANK charge added; see HELP RATES 399.
File 351:DERWENT WPI 1963-1998/UD=9819;UP=9816;UM=9814
(c)1998 Derwent Info Ltd
*File 351: Some images missing from UD=9816-9818 to be added as soon as
possible. Output formats changed for 1998. See HELP FORM 351 for info.

Set	Items	Description
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? s	(non(w)deplet? or nondeplet?) and (antibod? or immunoglobulin?)	

Processing

2777715	NON	
162261	DEPLET?	
270	NON(W)DEPLET?	
400	NONDEPLET?	
1034528	ANTIBOD?	
319454	IMMUNOGLOBULIN?	
S1	312	(NON(W)DEPLET? OR NONDEPLET?) AND (ANTIBOD? OR IMMUNOGLOBULIN?)

? s s1 and cd4

312	S1	
103565	CD4	
S2	224	S1 AND CD4

? s s2 and human?

Processing

224	S2	
9670595	HUMAN?	
S3	67	S2 AND HUMAN?

? rd s3

>>>Duplicate detection is not supported for File 351.

>>>Records from unsupported files will be retained in the RD set.
...examined 50 records (50)

...completed examining records
S4 45 RD S3 (unique items)
? t s4/3/all

4/3/1 (Item 1 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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14176107 BIOSIS Number: 01176107
Treatment of recalcitrant plaque psoriasis with a **humanized**
non-depleting antibody to CD4
Bachelez H; Flageul B; Dubertret L; Fraitag S; Grossman R; Brousse N;
Poisson D; Knowles R W; Wacholtz M C; Haverty T; Chatenoud L; Bach J-F
Hopital Necker, 161 Rue de Sevres, Paris, France
Journal of Autoimmunity 11 (1). 1998. 53-62.
Full Journal Title: Journal of Autoimmunity
ISSN: 0896-8411
Language: ENGLISH
Print Number: Biological Abstracts Vol. 105 Iss. 009 Ref. 118819

4/3/2 (Item 2 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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14157891 BIOSIS Number: 01157891

Reduction of Th1 cell activity in patients with rheumatoid arthritis after treatment with a **non-depleting** monoclonal **antibody** to **CD4**

Schulze-Koops H; Davis L S; Haverty P; Wacholtz M C; Lipsky P E
Southwestern Med. Cent., Dallas, TX, USA

Arthritis & Rheumatism 40 (9 SUPPL.). 1997. S191.

Full Journal Title: 61st National Scientific Meeting of the American College of Rheumatology and the 32nd National Scientific Meeting of the Association of Rheumatology Health Professionals, Washington, DC, USA, November 8-12, 1997. Arthritis & Rheumatism

ISSN: 0004-3591

Language: ENGLISH

Document Type: CONFERENCE PAPER

Print Number: Biological Abstracts/RRM Vol. 050 Iss. 004 Ref. 065775

4/3/3 (Item 3 from file: 55)

DIALOG(R) File 55:BIOSIS PREVIEWS(R)

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14157060 BIOSIS Number: 01157060

Effect of a **humanized non-depleting** anti-**CD4** monoclonal **antibody** (mAb) on synovial fluid (SF) in rheumatoid arthritis (RA)

Choy E H S; Connolly D J A; Rapson N; Kingsley G H; Johnston J M; Panayi G S

Rheumatol. Unit, Guy's and King's Coll. Hosp., London, UK

Arthritis & Rheumatism 40 (9 SUPPL.). 1997. S52.

Full Journal Title: 61st National Scientific Meeting of the American College of Rheumatology and the 32nd National Scientific Meeting of the Association of Rheumatology Health Professionals, Washington, DC, USA, November 8-12, 1997. Arthritis & Rheumatism

ISSN: 0004-3591

Language: ENGLISH

Document Type: CONFERENCE PAPER

Print Number: Biological Abstracts/RRM Vol. 050 Iss. 004 Ref. 064944

4/3/4 (Item 4 from file: 55)

DIALOG(R) File 55:BIOSIS PREVIEWS(R)

(c) 1998 BIOSIS. All rts. reserv.

14126123 BIOSIS Number: 01126123

Nondepleting humanized anti-**CD4** monoclonal **antibody** in patients with refractory rheumatoid arthritis

Moreland L W; Haverty T P; Wacholtz M C; Knowles R W; Bucy R P; Heck L W Jr; Koopman W J

Div. Rheumatology, Univ. Ala., 1717 6th Ave. South, Room 068, Birmingham, AL 35294-7201, USA

Journal of Rheumatology 25 (2). 1998. 221-228.

Full Journal Title: Journal of Rheumatology

ISSN: 0315-162X

Language: ENGLISH

Print Number: Biological Abstracts Vol. 105 Iss. 006 Ref. 084997

4/3/5 (Item 5 from file: 55)

DIALOG(R) File 55:BIOSIS PREVIEWS(R)

(c) 1998 BIOSIS. All rts. reserv.

14107479 BIOSIS Number: 01107479

Therapeutically effective **humanised non-depleting** anti-**CD4** monoclonal **antibody** (mAb) 4162W94 has no effect on monocyteoid cell lines

Newman I; Connolly D A; Choy E H S; Rapson N T; Panayi G S
Rheumatol. Unit, UMDS and King's Coll. Hosp., London SE1 9RT, UK
Immunology 92 (SUPPL. 1). 1997. 117.
Full Journal Title: 5th Annual Congress of the British Society for
Immunology, Brighton, England, UK, December 2-5, 1997. Immunology
ISSN: 0019-2805
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 050 Iss. 003 Ref. 041391

4/3/6 (Item 6 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

14010282 BIOSIS Number: 01010282
Humanized anti-**CD4** monoclonal **antibody** therapy of
autoimmune and inflammatory disease
Isaacs J D; Burrows N; Wing M; Keogan M T; Rebello P R U B; Watts R A;
Pye R J; Norris P; Hazelman B L; Hale G; Waldmann H
Molecular Med. Unit, Clin. Sci. Build., St. James's Univ. Hosp., Leeds
LS9 7TF, UK
Clinical and Experimental Immunology 110 (2). 1997. 158-166.
Full Journal Title: Clinical and Experimental Immunology
ISSN: 0009-9104
Language: ENGLISH
Print Number: Biological Abstracts Vol. 105 Iss. 001 Ref. 010282

4/3/7 (Item 7 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

13658881 BIOSIS Number: 99658881
A **humanized** form of a **CD4**-specific monoclonal **antibody**
exhibits decreased antigenicity and prolonged plasma half-life in rhesus
monkeys while retaining its unique biological and antiviral properties
Reimann K A; Lin W; Bixler S; Browning B; Ehrenfels B N; Lucci J;
Miatkowski K; Olson D; Parish T H; Rosa M D; Oleson F B; Hsu Y M; Padlan E
A; Letvin N L; Burkly L C
Division Viral Pathogenesis, Beth Israel Deaconess Med. Cent., RE-113,
330 Brookline Ave., Boston, MA 02215, USA
AIDS Research and Human Retroviruses 13 (11). 1997. 933-943.
Full Journal Title: AIDS Research and Human Retroviruses
ISSN: 0889-2229
Language: ENGLISH
Print Number: Biological Abstracts Vol. 104 Iss. 005 Ref. 067280

4/3/8 (Item 8 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

13627855 BIOSIS Number: 99627855
The immunological and pharmacodynamic effects of a **humanised**
non-depleting anti-**CD4** monoclonal **antibody** (mAb) in
rheumatoid arthritis (RA)
Choy E H S; Connolly D J A; Regan T; Manna V K; Rapson N; Kingsley G H;
Panayi G S; Johnston J M
Glaxo Wellcome, Beckenham, London, UK
British Journal of Rheumatology 36 (SUPPL. 1). 1997. 185.
Full Journal Title: XIVth Annual General Meeting of the British Society
of Rheumatology, Harrogate, England, UK, April 23-25, 1997. British
Journal of Rheumatology
ISSN: 0263-7103

Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 049 Iss. 008 Ref. 134420

4/3/9 (Item 9 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

13627731 BIOSIS Number: 99627731
The clinical effect of a by **humanised non-depleting**
anti-CD4 monoclonal antibody (mAb) in rheumatoid arthritis (RA)
Panayi G S; Choy E H S; Connolly D J A; Manna V K; Regan T; Rapson N;
Kingsley G H; Johnston J M
Rheumatology Unit, Guy's Hosp., UMDS, London, UK
British Journal of Rheumatology 36 (SUPPL. 1). 1997. 122.
Full Journal Title: XIVth Annual General Meeting of the British Society
of Rheumatology, Harrogate, England, UK, April 23-25, 1997. British
Journal of Rheumatology
ISSN: 0263-7103
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 049 Iss. 008 Ref. 134296

4/3/10 (Item 10 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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13402315 BIOSIS Number: 99402315
T cell hypothesis in rheumatoid arthritis (RA) tested by **humanised**
non-depleting anti-CD4 monoclonal antibody (mAb)
treatment I: Suppression of disease activity and acute phase response
Panayi G S; Choy E H S; Connolly D J A; Manna V K; Regan T; Rapson N;
Kingsley G H; Johnston J M
Rheumatology Unit, Guy's Hosp., UMDS, London, UK
Immunology 89 (SUPPL. 1). 1996. 92.
Full Journal Title: Joint Congress of the British Society for Immunology
and the Biochemical Society, Harrogate, England, UK, December 10-13, 1996.
Immunology
ISSN: 0019-2805
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 049 Iss. 003 Ref. 048954

4/3/11 (Item 11 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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13402314 BIOSIS Number: 99402314
T cell hypothesis in rheumatoid arthritis (RA) tested by **humanised**
non-depleting anti-CD4 monoclonal antibody (mAb)
treatment II: Clinical activity is related to pharmacodynamic effects
Choy E H S; Connolly D J A; Regan T; Manna V K; Rapson N; Kingsley G H;
Panayi G S; Johnston J M
Rheumatology Unit, Guy's Hosp., UMDS, London, UK
Immunology 89 (SUPPL. 1). 1996. 92.
Full Journal Title: Joint Congress of the British Society for Immunology
and the Biochemical Society, Harrogate, England, UK, December 10-13, 1996.
Immunology
ISSN: 0019-2805
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 049 Iss. 003 Ref. 048953

4/3/12 (Item 12 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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13224264 BIOSIS Number: 99224264
T cell hypothesis in rheumatoid arthritis (RA) tested by **humanized non-depleting anti-CD4 monoclonal antibody (mAb)**
treatment III: Immunological effects
Connolly D J A; Choy E H S; Rapson N; Regan T; Kingsley G H; Johnston J M
; Panayi G S
Rheumatol. Unit, Guy's Hosp., UMDS, London, UK
Arthritis & Rheumatism 39 (9 SUPPL.). 1996. S245.
Full Journal Title: 60th National Scientific Meeting of the American College of Rheumatology and the 31st National Scientific Meeting of the Association of Rheumatology Health Professionals, Orlando, Florida, USA, October 18-22, 1996. Arthritis & Rheumatism
ISSN: 0004-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 048 Iss. 011 Ref. 203385

4/3/13 (Item 13 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

13224263 BIOSIS Number: 99224263
T cell hypothesis in rheumatoid arthritis (RA) tested by **humanized non-depleting anti-CD4 monoclonal antibody (mAb)**
treatment II: Clinical activity is related to pharmacodynamic effects
Choy E H S; Connolly D J A; Regan T; Manna V K; Rapson N; Kingsley G H; Panayi G S; Johnston J M
Rheumatol. Unit, Guy's Hosp., UMDS, London, UK
Arthritis & Rheumatism 39 (9 SUPPL.). 1996. S244.
Full Journal Title: 60th National Scientific Meeting of the American College of Rheumatology and the 31st National Scientific Meeting of the Association of Rheumatology Health Professionals, Orlando, Florida, USA, October 18-22, 1996. Arthritis & Rheumatism
ISSN: 0004-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 048 Iss. 011 Ref. 203384

4/3/14 (Item 14 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

13224262 BIOSIS Number: 99224262
T cell hypothesis in rheumatoid arthritis (RA) tested by **humanized non-depleting anti-CD4 monoclonal antibody (mAb)**
treatment I: Suppression of disease activity and acute phase response
Panayi G S; Choy E H S; Connolly D J A; Regan T; Manna V K; Rapson N; Kingsley G H; Johnston J M
Rheumatol. Unit, Guy's Hosp., UMDS, London, UK
Arthritis & Rheumatism 39 (9 SUPPL.). 1996. S244.
Full Journal Title: 60th National Scientific Meeting of the American College of Rheumatology and the 31st National Scientific Meeting of the Association of Rheumatology Health Professionals, Orlando, Florida, USA, October 18-22, 1996. Arthritis & Rheumatism
ISSN: 0004-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER

4/3/15 (Item 15 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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13223537 BIOSIS Number: 99223537
Results of a placebo-controlled multicenter trial using a primatized
non-depleting, anti-**CD4** monoclonal **antibody** in the
treatment of rheumatoid arthritis
Levy R; Weisman M; Wisenhutter C; Yocum D; Schnitzer T; Goldman A; Schiff
M; Leiden B F; Solinger A; MacDonald B; Lipani J
Olympia, WA 98502, USA
Arthritis & Rheumatism 39 (9 SUPPL.). 1996. S122.
Full Journal Title: 60th National Scientific Meeting of the American
College of Rheumatology and the 31st National Scientific Meeting of the
Association of Rheumatology Health Professionals, Orlando, Florida, USA,
October 18-22, 1996. Arthritis & Rheumatism
ISSN: 0044-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 048 Iss. 011 Ref. 202658

4/3/16 (Item 16 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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13031632 BIOSIS Number: 99031632
Immunological markers of response in a multi-dose protocol 7002 using an
immunomodulating, **non-depleting** Primatized anti-**CD4**
monoclonal **antibody** in rheumatoid arthritis (RA)
Solinger A; Paxton H; Wey K; Yocum D
IDEC Pharmaceuticals, San Diego, CA 92121, USA
FASEB Journal 10 (6). 1996. A1314.
Full Journal Title: Joint Meeting of the American Society for
Biochemistry and Molecular Biology, the American Society for Investigative
Pathology and the American Association of Immunologists, New Orleans,
Louisiana, USA, June 2-6, 1996. FASEB Journal
ISSN: 0892-6638
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 048 Iss. 007 Ref. 125368

4/3/17 (Item 17 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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12165809 BIOSIS Number: 98765809
Immunological markers of response in a multi-dose protocol 7002 using an
immunomodulating, **non-depleting** primatized-TM anti-**CD4**
monoclonal **antibody** in rheumatoid arthritis (RA)
Solinger A; Paxton H; Wey K; Yocum D
IDEC Pharmaceuticals, San Diego, CA 92121, USA
FASEB Journal 10 (3). 1996. A442.
Full Journal Title: Experimental Biology 96, Part II, Washington, D.C.,
USA, April 14-17, 1996. FASEB Journal
ISSN: 0892-6638
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 048 Iss. 005 Ref. 082598

4/3/18 (Item 18 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

11935571 BIOSIS Number: 98535571
Modulation of mitogen and recall antigen proliferation by a **non-depleting**, anti-**CD4** monoclonal **antibody**: Results of a multi-dose study
Yocum D E; Mararescu M; Soundararaian D; Nordensson K; Solinger A M; Lipani J
Univ. Ariz., Tucson, AZ 85724, USA
Arthritis & Rheumatism 38 (9 SUPPL.). 1995. S280.
Full Journal Title: 59th National Scientific Meeting of the American College of Rheumatology and the 30th National Scientific Meeting of the Association of Rheumatology Health Professionals, San Francisco, California, USA, October 21-26, 1995. Arthritis & Rheumatism
ISSN: 0004-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 047 Iss. 012 Ref. 205446

4/3/19 (Item 19 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

11935007 BIOSIS Number: 98535007
Treating rheumatoid arthritis with a **non-depleting** anti-**CD4** monoclonal **antibody** (MAb)
Moreland L W; Bucy R P; Knowles R W; Wacholtz M C; Haverty T P; Koopman W J
Univ. Alabama at Birmingham, Birmingham, AL, USA
Arthritis & Rheumatism 38 (9 SUPPL.). 1995. S186.
Full Journal Title: 59th National Scientific Meeting of the American College of Rheumatology and the 30th National Scientific Meeting of the Association of Rheumatology Health Professionals, San Francisco, California, USA, October 21-26, 1995. Arthritis & Rheumatism
ISSN: 0004-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 047 Iss. 012 Ref. 204882

4/3/20 (Item 20 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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11935003 BIOSIS Number: 98535003
Results of a multi-dose protocol 7002 using an immunomodulating, **non-depleting** PRIMATIZED anti-**CD4** monoclonal **antibody** in rheumatoid arthritis (RA)
Kaine J; Solinger A; Yocum D; Lipani J; Klas P; Tesser J; Wiesenhutter C; O'Sullivan F; Shuman S; Rigby W
Sarasota Arthritis Center, Sarasota, FL 34239, USA
Arthritis & Rheumatism 38 (9 SUPPL.). 1995. S185.
Full Journal Title: 59th National Scientific Meeting of the American College of Rheumatology and the 30th National Scientific Meeting of the Association of Rheumatology Health Professionals, San Francisco, California, USA, October 21-26, 1995. Arthritis & Rheumatism
ISSN: 0004-3591
Language: ENGLISH
Document Type: CONFERENCE PAPER
Print Number: Biological Abstracts/RRM Vol. 047 Iss. 012 Ref. 204878

4/3/21 (Item 21 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

11922318 BIOSIS Number: 98522318
Therapeutic monoclonal **antibodies**
Choy E H S; Panayi G S; Kingsley G H
Rheumatol. Unit, Div. Medicine, UMDS, 4th Floor, Hunt's House, Guy's
Hospital, St. Thomas Street, London SE1 9RT, UK
British Journal of Rheumatology 34 (8). 1995. 707-715.
Full Journal Title: British Journal of Rheumatology
ISSN: 0263-7103
Language: ENGLISH
Print Number: Biological Abstracts Vol. 100 Iss. 011 Ref. 173175

4/3/22 (Item 22 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

11760328 BIOSIS Number: 98360328
Activation of CD4+ T cells in the presence of a **nondepleting**
monoclonal **antibody** to CD4 induces a Th2-Type response in vitro
Stumbles P; Mason D
MRC Cellular Immunol. Unit, Sir William Dunn Sch. Pathol., University
Oxford, South Parks Rd., Oxford OX1 3RE, UK
Journal of Experimental Medicine 182 (1). 1995. 5-13.
Full Journal Title: Journal of Experimental Medicine
ISSN: 0022-1007
Language: ENGLISH
Print Number: Biological Abstracts Vol. 100 Iss. 004 Ref. 052166

4/3/23 (Item 23 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

11345669 BIOSIS Number: 97545669
Immunological approach to inhibit formation of anti-**antibodies** to
allo- and xenogeneic anti-T cell **immunoglobulin**
Mysliwietz J; Thierfelder S; Mocikat R; Kremmer E
GSF, Inst. Immunol., Marchioninistr. 25, D-81377 Muenchen, GER
European Journal of Immunology 24 (10). 1994. 2323-2328.
Full Journal Title: European Journal of Immunology
ISSN: 0014-2980
Language: ENGLISH
Print Number: Biological Abstracts Vol. 098 Iss. 012 Ref. 163292

4/3/24 (Item 24 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

10805769 BIOSIS Number: 97005769
T-cell recognition of a cross-reactive antigen(s) in erythrocytic stages
of Plasmodium falciparum and Plasmodium yoelii: Inhibition of parasitemia
by this antigen(s)
Lucas B; Engels A; Camus D; Haque A
Centre Immunol., Biol. Parasitaire, Inst. Pasteur, 59019 Lille, FRA
Infection and Immunity 61 (11). 1993. 4863-4869.
Full Journal Title: Infection and Immunity
ISSN: 0019-9567
Language: ENGLISH
Print Number: Biological Abstracts Vol. 097 Iss. 001 Ref. 005267

4/3/25 (Item 25 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

8095291 BIOSIS Number: 91016291
RESISTANCE TO INFECTION BY HIV-1 OF PERIPHERAL BLOOD MONONUCLEAR CELLS
FROM HIV-1-INFECTED PATIENTS IS PROBABLY MEDIATED BY NEUTRALIZING
ANTIBODIES

TREMBLAY M; NUMAZAKI K; LI X; GORNITSKY M; HISCOTT J; WAINBERG M A
MCGILL AIDS CENTRE JEWISH GENERAL HOSP., 3755 COTE STE-CATHERINE ROAD,
MONTREAL, QUEBEC H3T 1E2, CAN.
J IMMUNOL 145 (9). 1990. 2896-2901. CODEN: JOIMA
Full Journal Title: Journal of Immunology
Language: ENGLISH

4/3/26 (Item 1 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

10623307 EMBASE No: 98050169
Clinical pharmacology and therapeutic potential of monoclonal
antibody treatment in rheumatoid arthritis
Choy E.H.S.
Dr. E.H.S. Choy, Rheumatology Unit, Thomas Guy House, Guy's Hospital, St
Thomas Street, London SE1 9RT United Kingdom
Drugs and Aging (New Zealand) , 1998, 12/2 (139-148)
CODEN: DRAGE ISSN: 1170-229X
DOCUMENT TYPE: Journal Review
LANGUAGES: ENGLISH SUMMARY LANGUAGES: ENGLISH
NUMBER OF REFERENCES: 51

4/3/27 (Item 2 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

9787616 EMBASE No: 95351540
T-cell regulation
Choy E.H.S.; Kingsley G.H.; Panayi G.S.
UMDS, Rheumatology Unit, Guy's Hospital, St Thomas Street, London SE1 9RT
United Kingdom
Bailliere's Clinical Rheumatology (United Kingdom) , 1995, 9/4 (653-671)
CODEN: BCRHE ISSN: 0950-3579
LANGUAGES: English SUMMARY LANGUAGES: English

4/3/28 (Item 3 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

9532547 EMBASE No: 95106020
Anti-**CD4** monoclonal **antibody** immune intervention in patients
with newly diagnosed Type I (insulin-dependent) diabetes mellitus
Hehmke B.; Kuttler B.; Laube F.; Gens E.; Michaelis D.; Hahn H.-J.;
Schulze-Koops H.; Emmrich F.
Institute Diabetes 'Gerhardt Katsch', Dept Experimental Clin
Endocrinology, D-17495 Karlsburg Germany
Diabetes, Nutrition and Metabolism - Clinical and Experimental (Italy) ,
1994, 7/5 (273-280)
CODEN: DNMEE ISSN: 0394-3402
LANGUAGES: English SUMMARY LANGUAGES: English

4/3/29 (Item 4 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

8675097 EMBASE No: 92355607
Anti-**CD4** monoclonal **antibodies** in therapy: Creation of nonclassical tolerance in the adult
Shizuru J.A.; Alters S.E.; Fathman C.G.
Stanford Univ. School of Medicine, Div. of Rheumatology and Immunology, Stanford, CA 94305 USA
IMMUNOL. REV. (Denmark) , 1992, -/129 (105-130)
CODEN: IMRED ISSN: 0105-2896 ADONIS ORDER NUMBER: 010528969200046X
LANGUAGES: English SUMMARY LANGUAGES: English

4/3/30 (Item 5 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

8398766 EMBASE No: 92074758
Comparison of GK1.5 and chimeric rat/mouse GK1.5 anti-**CD4** **antibodies** for prolongation of skin allograft survival and suppression of alloantibody production in mice
Rashid A.; Auchincloss H. Jr.; Sharon J.
Boston University School of Medicine, 80 East Concord St., Boston, MA 02118 USA
J. IMMUNOL. (USA) , 1992, 148/5 (1382-1388)
CODEN: JOIMA ISSN: 0022-1767
LANGUAGES: English SUMMARY LANGUAGES: English

4/3/31 (Item 6 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

8183556 EMBASE No: 91209639
Monoclonal **antibody** therapy for the induction of transplantation tolerance
Cobbold S.P.
Division of Immunology, Cambridge University Department of Pathology, Tennis Court Road, Cambridge CB1 2QP United Kingdom
IMMUNOL. LETT. (Netherlands) , 1991, 29/1-2 (117-122)
CODEN: IMLED ISSN: 0165-2478 ADONIS ORDER NUMBER: 016524789100175N
LANGUAGES: English

4/3/32 (Item 7 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

8013038 EMBASE No: 91038466
Induction of tolerance in peripheral T cells with monoclonal **antibodies**
Qin S.; Wise M.; Cobbold S.P.; Leong L.; Kong Y.-C.M.; Parnes J.R.; Waldmann H.
Division of Immunology, Department of Pathology, Cambridge University, Cambridge CB2 2QQ United Kingdom
EUR. J. IMMUNOL. (Germany, Federal Republic of) , 1990, 20/12 (2737-2745)
CODEN: EJIMA ISSN: 0014-2980
LANGUAGES: English

4/3/33 (Item 1 from file: 154)
DIALOG(R) File 154:MEDLINE(R)
(c) format only 1998 Dialog Corporation. All rts. reserv.

09479687 98184497

Mucosal immunity to herpes simplex virus type 2 infection in the mouse vagina is impaired by in vivo depletion of T lymphocytes.

Parr MB; Parr EL

School of Medicine, Southern Illinois University, Carbondale 62901, USA.
mparr@som.siu.edu

J Virol (UNITED STATES) Apr 1998, 72 (4) p2677-85, ISSN 0022-538X
Journal Code: KCV

Contract/Grant No.: HD-17337, HD, NICHD

Languages: ENGLISH

Document type: JOURNAL ARTICLE

4/3/34 (Item 2 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

(c) format only 1998 Dialog Corporation. All rts. reserv.

08916757 97172248

Induction of donor specific transplantation tolerance to cardiac allografts following treatment with **nondepleting** (RIB 5/2) or depleting (OX-38) anti-**CD4** mAb plus intrathymic or intravenous donor alloantigen.

Arima T; Lehmann M; Flye MW

Department of Surgery, Washington University School of Medicine, St. Louis, Missouri, USA.

Transplantation (UNITED STATES) Jan 27 1997, 63 (2) p284-92, ISSN 0041-1337 Journal Code: WEJ

Contract/Grant No.: 5PO1 AI35121, AI, NIAID

Languages: ENGLISH

Document type: JOURNAL ARTICLE

4/3/35 (Item 3 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

(c) format only 1998 Dialog Corporation. All rts. reserv.

08909043 97111516

Nondepleting anti-**CD4** antibody treatment prolongs lung-directed El-deleted adenovirus-mediated gene expression in rats.

Lei D; Lehmann M; Shellito JE; Nelson S; Siegling A; Volk HD; Kolls JK
LSU Section of Pulmonary/Critical Care MEB, New Orleans 70112, USA.

Hum Gene Ther (UNITED STATES) Dec 1 1996, 7 (18) p2273-9, ISSN 1043-0342 Journal Code: A12

Contract/Grant No.: R29-AA10384, AA, NIAAA; HL-29246, HL, NHLBI

Languages: ENGLISH

Document type: JOURNAL ARTICLE

4/3/36 (Item 4 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

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08548769 96161423

Innovative treatment approaches for rheumatoid arthritis. T-cell regulation.

Choy EH; Kingsley GH; Panayi GS

UMDS, Rheumatology Unit, Guy's Hospital, London, UK.

Baillieres Clin Rheumatol (ENGLAND) Nov 1995, 9 (4) p653-71, ISSN 0950-3579 Journal Code: CRY

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

4/3/37 (Item 5 from file: 154)

DIALOG(R) File 154: MEDLINE(R)
(c) format only 1998 Dialog Corporation. All rts. reserv.

08010998 94321135
Sparing of the ipsilateral retina after anterior chamber inoculation of HSV-1: requirement for either CD4+ or CD8+ T cells.
Azumi A; Atherton SS
Department of Cellular and Structural Biology, University of Texas Health Science Center, San Antonio 78284-7762.
Invest Ophthalmol Vis Sci (UNITED STATES) Jul 1994, 35 (8) p3251-9,
ISSN 0146-0404 Journal Code: GWI
Contract/Grant No.: EY06012, EY, NEI
Languages: ENGLISH
Document type: JOURNAL ARTICLE

4/3/38 (Item 6 from file: 154)
DIALOG(R) File 154: MEDLINE(R)
(c) format only 1998 Dialog Corporation. All rts. reserv.

08005292 94131763
Modulation of murine herpes simplex virus type 1 retinitis in the uninoculated eye by CD4+ T lymphocytes.
Azumi A; Cousins SW; Kanter MY; Atherton SS
Department of Cellular and Structural Biology, University of Texas Health Science Center, San Antonio 78284-7762.
Invest Ophthalmol Vis Sci (UNITED STATES) Jan 1994, 35 (1) p54-63,
ISSN 0146-0404 Journal Code: GWI
Contract/Grant No.: EY06012, EY, NEI
Languages: ENGLISH
Document type: JOURNAL ARTICLE

4/3/39 (Item 7 from file: 154)
DIALOG(R) File 154: MEDLINE(R)
(c) format only 1998 Dialog Corporation. All rts. reserv.

07351119 93153366
Down regulation of stem cell colony formation by purified CD8 lymphocytes and CD8 conditioned medium: potential importance for bone marrow transplantation in leukemia.
Gazitt Y; He YJ
Department of Pediatric Hematology-Oncology, University of Florida, Gainesville.
Leuk Lymphoma (SWITZERLAND) Sep 1992, 8 (1-2) p117-27, ISSN 1042-8194
Journal Code: BNQ
Languages: ENGLISH
Document type: JOURNAL ARTICLE

4/3/40 (Item 8 from file: 154)
DIALOG(R) File 154: MEDLINE(R)
(c) format only 1998 Dialog Corporation. All rts. reserv.

07342782 92368404
The forces driving autoimmune disease.
Roitt IM; Hutchings PR; Dawe KI; Sumar N; Bodman KB; Cooke A
Dept. of Immunology, University College & Middlesex School of Medicine, London, UK.
J Autoimmun (ENGLAND) Apr 1992, 5 Suppl A p11-26, ISSN 0896-8411
Journal Code: ADL
Languages: ENGLISH
Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

4/3/41 (Item 9 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

(c) format only 1998 Dialog Corporation. All rts. reserv.

06603039 91370929

Reprogramming the immune system for tolerance with monoclonal
antibodies.

Cobbold SP; Qin SX; Waldmann H

Department of Pathology, Cambridge University, UK.

Semin Immunol (UNITED STATES) Nov 1990, 2 (6) p377-87, ISSN 1044-5323

Journal Code: A61

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

4/3/42 (Item 10 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

(c) format only 1998 Dialog Corporation. All rts. reserv.

05517371 89001400

CD4+ T cells appear capable of initiating graft-versus-host disease across non-major histocompatibility complex (MHC) barriers in man.

Atkinson K; Cooley M; Farrelly H; O'Flaherty E; Ashby M; Biggs J

Department of Haematology, St Vincent's Hospital, Sydney, Australia.

Bone Marrow Transplant (ENGLAND) Jun 1987, 2 (1) p79-84, ISSN

0268-3369 Journal Code: BON

Languages: ENGLISH

Document type: JOURNAL ARTICLE

4/3/43 (Item 1 from file: 351)

DIALOG(R) File 351: DERWENT WPI

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011033929

WPI Acc No: 97-011853/199701

XRAM Acc No: C97-003237

Amt. of **non-depleting** anti-**CD4** antibody effective

to induce immunological tolerance - useful to inhibit allo-graft rejection in primate subject, specifically bone marrow allo-graft

Patent Assignee: JOHNSON & JOHNSON CORP (JOHJ)

Inventor: CAVENDER D E; KNOWLES R W; THOMAS J M

Number of Countries: 069 Number of Patents: 002

Patent Family:

Patent No	Kind	Date	Applicat No	Kind	Date	Main IPC	Week
WO 9636359	A1	19961121	WO 96US6912	A	19960516	A61K-039/395	199701 B
AU 9657479	A	19961129	AU 9657479	A	19960516	A61K-039/395	199712

Priority Applications (No Type Date): US 95443739 A 19950518

Filing Details:

Patent	Kind	Filing Notes	Application	Patent
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WO 9636359 A1

Designated States (National): AL AM AT AU AZ BB BG BR BY CA CH CN CZ DE DK EE ES FI GB GE HU IS JP KE KG KP KR KZ LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG UZ VN

Designated States (Regional): AT BE CH DE DK EA ES FI FR GB GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG

AU 9657479 A Based on

WO 9636359

Language, Pages: WO 9636359 (E, 17)

4/3/44 (Item 2 from file: 351)

DIALOG(R) File 351: DERWENT WPI

(c) 1998 Derwent Info Ltd. All rts. reserv.

009140953

WPI Acc No: 92-268391/199232

XRAM Acc No: C92-119699

Use of single **non-depleting CD4** monoclonal antibody - for treatment of insulin-dependent diabetes mellitus (IDDM), arrests loss of insulin producing cells

Patent Assignee: UNIV COLLEGE LONDON (UNLO); GLAXO WELLCOME PLC (GLAX)

Inventor: COOKE A; WALDMANN H

Number of Countries: 035 Number of Patents: 009

Patent Family:

Patent No	Kind	Date	Applicant	No	Kind	Date	Main IPC	Week
WO 9211869	A1	19920723	WO	92GB74	A	19920114	A61K-039/395	199232 B
AU 9211647	A	19920817	AU	9211647	A	19920114	A61K-039/395	199245
			WO	92GB74	A	19920114		
EP 567490	A1	19931103	EP	92902288	A	19920114	A61K-039/395	199344
			WO	92GB74	A	19920114		
JP 6504283	W	19940519	JP	92502777	A	19920114	A61K-039/395	199424
			WO	92GB74	A	19920114		
AU 668081	B	19960426	AU	9211647	A	19920114	A61K-039/395	199624
EP 567490	B1	19970813	EP	92902288	A	19920114	A61K-039/395	199737
			WO	92GB74	A	19920114		
DE 69221605	E	19970918	DE	621605	A	19920114	A61K-039/395	199743
			EP	92902288	A	19920114		
			WO	92GB74	A	19920114		
US 5670150	A	19970923	US	9390203	A	19931201	A61K-039/395	199744
			US	95436843	A	19950508		
ES 2106169	T3	19971101	EP	92902288	A	19920114	A61K-039/395	199750

Priority Applications (No Type Date): GB 91741 A 19910114

Filing Details:

Patent	Kind	Filing Notes	Application	Patent
WO 9211869	A1			
		Designated States (National): AT AU BB BG BR CA CH DE DK ES FI GB HU JP KP KR LK LU MG MW NL NO PL RO RU SD SE US		
		Designated States (Regional): AT BE CH DE DK ES FR GB GR IT LU MC NL OA SE		
AU 9211647	A	Based on		WO 9211869
EP 567490	A1	Based on		WO 9211869
		Designated States (Regional): AT BE CH DE DK ES FR GB GR IT LI LU MC NL SE		
JP 6504283	W	Based on		WO 9211869
AU 668081	B	Previous Publ.		AU 9211647
		Based on		WO 9211869
EP 567490	B1	Based on		WO 9211869
		Designated States (Regional): AT BE CH DE DK ES FR GB GR IT LI LU MC NL SE		
DE 69221605	E	Based on		EP 567490
		Based on		WO 9211869
US 5670150	A	Cont of	US 9390203	
ES 2106169	T3	Based on		EP 567490
Language, Pages: WO 9211869 (E, 19); EP 567490 (E); JP 6504283 (5); EP 567490 (E, 6); US 5670150 (5)				

4/3/45 (Item 3 from file: 351)

DIALOG(R) File 351:DERWENT WPI

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008503137

WPI Acc No: 91-007221/199101

XRAM Acc No: C91-003203

Non-depleting CD4 and **CD8** monoclonal **antibodies** -
for inducing tolerance to foreign antigens in transplant rejection,
auto-immune disease, etc

Patent Assignee: COBBOLD S P (COBB-I); WALDMANN H (WALD-I); WELLCOME FOUND

LTD (WELL); GLAXO WELLCOME INC (GLAX)
 Inventor: COBBOLD S P; WALDMANN H
 Number of Countries: 025 Number of Patents: 015
 Patent Family:

Patent No	Kind	Date	Applicat No	Kind	Date	Main IPC	Week
WO 9015152	A	19901213					199101 B
PT 94214	A	19910208					199109
AU 9057258	A	19910107					199115
EP 474691	A	19920318	EP 90908270	A	19900531		199212
ZA 9004174	A	19920226	ZA 904174	A	19900530		199213
DD 296843	A5	19911219	DD 341218	A	19900531	A61K-039/395	199221
JP 4505919	W	19921015	JP 90508030	A	19900531	A61K-039/395	199248
			WO 90GB840	A	19900531		
HU 61341	T	19921230	HU 905134	A	19900531	C12P-021/08	199306
			WO 90GB840	A	19900531		
AU 657255	B	19950309	AU 9057258	A	19900531	C12P-021/08	199520
EP 474691	B1	19961113	EP 90908270	A	19900531	C12P-021/08	199650
			WO 90GB840	A	19900531		
DE 69029134	E	19961219	DE 629134	A	19900531	C12P-021/08	199705
			EP 90908270	A	19900531		
			WO 90GB840	A	19900531		
ES 2096588	T3	19970316	EP 90908270	A	19900531	C12P-021/08	199718
NZ 233889	A	19970624	NZ 233889	A	19900531	A61K-039/395	199732
BR 1100287	A3	19970916	BR 971100287	A	19970415	C12P-021/08	199744
US 5690933	A	19971125	US 91768868	A	19910727	A61K-039/395	199802
			US 9347344	A	19930329		
			US 94181170	A	19940113		
			US 94289532	A	19940812		

Priority Applications (No Type Date): GB 8912497 A 19890531

Filing Details:

Patent	Kind	Filing Notes	Application	Patent
WO 9015152	A	Designated States (National): AU CA FI HU JP KR US		
		Designated States (Regional): AT BE CH DE DK ES FR GB IT LU NL SE		
EP 474691	A	Designated States (Regional): AT BE CH DE DK ES FR GB IT LI LU NL SE		
JP 4505919	W	Based on	WO 9015152	
HU 61341	T	Based on	WO 9015152	
AU 657255	B	Previous Publ.	AU 9057258	
		Based on	WO 9015152	
EP 474691	B1	Based on	WO 9015152	
		Designated States (Regional): AT BE CH DE DK ES FR GB IT LI LU NL SE		
DE 69029134	E	Based on	EP 474691	
		Based on	WO 9015152	
ES 2096588	T3	Based on	EP 474691	
US 5690933	A	Cont of	US 91768868	
		Cont of	US 9347344	
		Cont of	US 94181170	

Language, Pages: EP 474691 (44); ZA 9004174 (57); JP 4505919 (19); EP 474691 (E, 32); US 5690933 (23)

? ds

Set	Items	Description
S1	312	(NON(W)DEPLET? OR NONDEPLET?) AND (ANTIBOD? OR IMMUNOGLOBULIN?)
S2	224	S1 AND CD4
S3	67	S2 AND HUMAN?
S4	45	RD S3 (unique items)

? s s1 and cd8

312	S1	
61088	CD8	
S5	95	S1 AND CD8

? rd s5

>>>Duplicate detection is not supported for File 351.

>>>Records from unsupported files will be retained in the RD set.

...examined 50 records (50)

...completed examining records

 S6 42 RD S5 (unique items)

? s s2 and review?

 224 S2

 2149145 REVIEW?

 S7 6 S2 AND REVIEW?

? t s7/3/all

7/3/1 (Item 1 from file: 55)

DIALOG(R) File 55:BIOSIS PREVIEWS(R)

(c) 1998 BIOSIS. All rts. reserv.

11922318 BIOSIS Number: 98522318

Therapeutic monoclonal **antibodies**

Choy E H S; Panayi G S; Kingsley G H

Rheumatol. Unit, Div. Medicine, UMDS, 4th Floor, Hunt's House, Guy's Hospital, St. Thomas Street, London SE1 9RT, UK

British Journal of Rheumatology 34 (8). 1995. 707-715.

Full Journal Title: British Journal of Rheumatology

ISSN: 0263-7103

Language: ENGLISH

Print Number: Biological Abstracts Vol. 100 Iss. 011 Ref. 173175

7/3/2 (Item 1 from file: 72)

DIALOG(R) File 72:EMBASE

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10623307 EMBASE No: 98050169

Clinical pharmacology and therapeutic potential of monoclonal **antibody** treatment in rheumatoid arthritis

Choy E.H.S.

Dr. E.H.S. Choy, Rheumatology Unit, Thomas Guy House, Guy's Hospital, St Thomas Street, London SE1 9RT United Kingdom

Drugs and Aging (New Zealand) , 1998, 12/2 (139-148)

CODEN: DRAGE ISSN: 1170-229X

DOCUMENT TYPE: Journal Review

LANGUAGES: ENGLISH SUMMARY LANGUAGES: ENGLISH

NUMBER OF REFERENCES: 51

7/3/3 (Item 2 from file: 72)

DIALOG(R) File 72:EMBASE

(c) 1998 Elsevier Science B.V. All rts. reserv.

9787616 EMBASE No: 95351540

T-cell regulation

Choy E.H.S.; Kingsley G.H.; Panayi G.S.

UMDS, Rheumatology Unit, Guy's Hospital, St Thomas Street, London SE1 9RT United Kingdom

Bailliere's Clinical Rheumatology (United Kingdom) , 1995, 9/4 (653-671)

CODEN: BCRHE ISSN: 0950-3579

LANGUAGES: English SUMMARY LANGUAGES: English

7/3/4 (Item 3 from file: 72)

DIALOG(R) File 72:EMBASE

9737958 EMBASE No: 95293479
Therapeutic monoclonal **antibodies**
Choy E.H.S.; Panayi G.S.; Kingsley G.H.
Rheumatology Unit, Division of Medicine, UMDS, Guy's Hospital, St Thomas
Street, London SE1 9RT United Kingdom
British Journal of Rheumatology (United Kingdom) , 1995, 34/8 (707-715)
CODEN: BJRHD ISSN: 0263-7103
LANGUAGES: English SUMMARY LANGUAGES: English

7/3/5 (Item 4 from file: 72)
DIALOG(R) File 72:EMBASE
(c) 1998 Elsevier Science B.V. All rts. reserv.

8675097 EMBASE No: 92355607
Anti-**CD4** monoclonal **antibodies** in therapy: Creation of
nonclassical tolerance in the adult
Shizuru J.A.; Alters S.E.; Fathman C.G.
Stanford Univ. School of Medicine, Div. of Rheumatology and Immunology,
Stanford, CA 94305 USA
IMMUNOL. REV. (Denmark) , 1992, -/129 (105-130)
CODEN: IMRED ISSN: 0105-2896 ADONIS ORDER NUMBER: 010528969200046X
LANGUAGES: English SUMMARY LANGUAGES: English

7/3/6 (Item 1 from file: 399)
DIALOG(R) File 399:CA SEARCH(R)
(c) 1998 American Chemical Society. All rts. reserv.

128191337 CA: 128(16)191337j CONFERENCE PROCEEDING
The therapeutic potential of a primatized nondepleting anti-CD4
(IDEC-CE9.1) monoclonal antibody in rheumatoid arthritis
AUTHOR(S): Solinger, Alan M.; Truneh, Alemseged; Lipani, John A.; Newman,
Roland A.
LOCATION: IDEC Pharmaceutical Corporation, San Diego, CA, USA
JOURNAL: Antibody Ther. EDITOR: Harris, William J. (Ed), Adair, John R
(Ed), DATE: 1997 PAGES: 341-353 CODEN: 65RLAP LANGUAGE: English
PUBLISHER: CRC, Boca Raton, Fla
? s s6 and human?

Processing

42 S6
9670595 HUMAN?
S8 10 S6 AND HUMAN?
? rd s8

>>>Duplicate detection is not supported for File 351.

>>>Records from unsupported files will be retained in the RD set.
...completed examining records
S9 10 RD S8 (unique items)
? t s9/7/all

9/7/1 (Item 1 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

11345669 BIOSIS Number: 97545669
Immunological approach to inhibit formation of anti-**antibodies** to
allo- and xenogeneic anti-T cell **immunoglobulin**
Mysliwietz J; Thierfelder S; Mocikat R; Kremmer E
GSF, Inst. Immunol., Marchioninistr. 25, D-81377 Muenchen, GER

Inhibitory anti-**antibodies** induced in patients by xenogeneic or even by **humanized** anti-T cell **antibodies** remain an unresolved problem. Mice also produce anti-**antibodies** following injection of xeno- or allogeneic anti-T cell **antibodies**. Here we report a principle based on sequentially applied anti-T cell **antibodies** generated in different species, which results in suppressed anti-**antibody** formation and prolonged immunosuppression. Thus, a single priming injection in mice of mouse (MmT1 or MmT5 differing by idioype only) or of rat (RmT1) anti-mouse Thy-1 monoclonal **antibodies** (mAb) or of rat anti-mouse L3T4 + Ly-2 (RmCD4 + **CD8**) mAb suppressed anti-**antibody** formation against subsequent booster injections of one of the above **antibodies**, provided that they differed in species origin from the priming **antibody**. Correspondingly, a sixfold and longer prolongation of 50 % survival of fully mismatched skin grafts was observed. Less or no anti-**antibody** suppression and little prolongation of graft survival was obtained if the 'first' and the 'second' (and following) **antibody** injections were of the same species, differing by iso- or idioype only. Finally, the suppressive principle did not manifest itself at all if the initial **antibody** injection included both the first and second **antibody**. These findings are discussed with reference to earlier studies on hapten/carrier effects as well as on immunosuppression attributed to 'non-depleting' rat anti-CD4/**CD8** T cell **antibodies**.

9/7/2 (Item 2 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

10805769 BIOSIS Number: 97005769

T-cell recognition of a cross-reactive antigen(s) in erythrocytic stages of Plasmodium falciparum and Plasmodium yoelii: Inhibition of parasitemia by this antigen(s)

Lucas B; Engels A; Camus D; Haque A

Centre Immunol., Biol. Parasitaire, Inst. Pasteur, 59019 Lille, FRA
Infection and Immunity 61 (11). 1993. 4863-4869.

Full Journal Title: Infection and Immunity

ISSN: 0019-9567

Language: ENGLISH

Print Number: Biological Abstracts Vol. 097 Iss. 001 Ref. 005267

In the current study, we investigated the presence of a cross-reactive antigen(s) in the erythrocyte stage from Plasmodium yoelii (265 BY strain) and Plasmodium falciparum through recognition by T cells primed in vivo with antigens from each of these parasites. BALB/c mice are naturally resistant to P. falciparum but are susceptible to P. yoelii infection. Mice that had recovered from P. yoelii primary infection became resistant to a second infection. A higher in vitro proliferative response to a soluble blood stage preparation of P. falciparum was observed in splenic cells from immune animals than in those from mice with a patent P. yoelii infection. The antigen-induced proliferative response was enhanced when animals were exposed to a secondary infection. Animals exposed to a challenge infection were treated with anti-CD4 or anti-**CD8** monoclonal **antibodies** to deplete the corresponding subset of T cells. There was a marked diminution in P. falciparum antigen-induced proliferative response in the total splenic cell populations from **CD8**-depleted but not from CD4-depleted mice. In **CD8**-depleted and **nondepleted** animals, the antigen-induced proliferation in the total cell populations was markedly lower than in the T-cell-rich populations, indicating inhibitory activities of B cells and/or macrophages. There was no such difference in the stimulation between total and T-enriched cell populations from CD4-depleted

animals. Flow cytometry analysis demonstrated the presence of an almost equal percentage of **CD8+** (59.6%) and CD4+ (64%) T cells in the spleen preparations following in vivo depletion of CD4- and **CD8**-bearing T cells, respectively. When cultured with *P. yoelii* blood stage antigen, splenocytes from animals immunized with *P. falciparum* antigen displayed a significant proliferative response which was markedly diminished by treatment with anti-Thy-1.2 **antibody** plus complement. Animals immunized with *P. falciparum* antigen and then challenged with *P. yoelii* blood stage parasites displayed about a 50% lower level of parasitemia. These results demonstrated the existence of a cross-reactive antigen(s) between a murine and a **human** *Plasmodium* species, as determined from both in vivo and in vitro biological assays, and indicated the reactivity of mainly **CD8+** T cells with this antigen.

9/7/3 (Item 3 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

7083977 BIOSIS Number: 88006722
ENDOGENOUSLY GENERATED ACTIVATED KILLER CELLS CIRCULATE AFTER AUTOLOGOUS AND ALLOGENEIC MARROW TRANSPLANTATION BUT NOT AFTER CHEMOTHERAPY
REITTIE J E; GOTTLIEB D; HESLOP H E; LEGER O; DEXLER H G; HAZLEHURST G; HOFFBRAND A V; PRENTICE H G; BRENNER M K

DEP. HAEMATOL., ROYAL FREE HOSP., POND ST., LONDON, NW3, UK.
BLOOD 73 (5). 1989. 1351-1358. CODEN: BLOOA

Full Journal Title: Blood

Language: ENGLISH

After marrow transplantation, major histocompatibility complex (MHC)-unrestricted natural killer (NK) lymphocytes are among the first cells to appear in the circulation. After T-cell-depleted bone marrow transplantation (TD-BMT), these cells have an activated pattern of target cell killing; they also secrete lymphokines including .gamma.-interferon (.gamma.-IFN), interleukin-2 (IL-2), and tumor necrosis factor (TNF) and may have a significant role as a primary defense against viral reactivation and in the elimination of residual host malignancy. We studied 43 patients with hematologic malignancy, treated by allogeneic TD-BMT, autologous **nondepleted** BMT, or chemotherapy alone to investigate (a) the mechanisms underlying the generation of these activated killer cells, (b) the range of conditions under which they are produced, and (c) their surface phenotype. We showed that .gamma.-IFN-secreting activated killer cells with the capacity to kill MHC-nonidentical NK-resistant targets are generated 4 to 6 weeks after either allogeneic TD-BMT or autologous BMT but do not appear after treatment with chemotherapy. Production therefore is not owing to T-cell depletion per se or to host donor alloreactivity, nor is it caused by stimulation by alloantigens contained in blood product support since no significant difference exists between allograft and chemotherapy patients in the number of units of blood platelet support given in the posttreatment period. Because most patients had no evidence of stimulation from virus reactivation/infection, the phenomenon of activation therefore appears to represent posttransplant immune disregulation following repopulation of the host immune system with lymphoid subsets derived exclusively from blood and marrow. Activated killing is predominantly mediated by the CD16+ CD3- subset, but substantial activity remains in the CD16- CD3+ cell fraction. Monoclonal **antibodies** (MoAbs) that block interaction with class-I MHC molecules at the level of target cell (W6/32 anti-HLA class I) or effector cell (**CD8**) do not inhibit killing by CD16- CD3+ cells. Activated killer cells may contribute to the lower risk of relapse after marrow transplantation as compared with intensive chemotherapy.

9/7/4 (Item 4 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

5856041 BIOSIS Number: 83118348

A COMPARATIVE STUDY OF T-CELL DEPLETED AND NON-DEPLETED MARROW TRANSPLANTATION FOR HEMATOLOGICAL MALIGNANCY

ATKINSON K; ASHBY M; BIGGS J; CONCANNON A; COOLEY M; DODDS A; FARRELLY H; MORGAN G; O'FLAHERTY E; ET AL

BONE MARROW TRANSPLANT. UNIT, ST. VICENT'S HOSP., DARLINGHURST, NSW 2010. AUST N Z J MED 17 (1). 1987. 16-23. CODEN: ANZJB

Full Journal Title: Australian and New Zealand Journal of Medicine

Language: ENGLISH

Sixteen patients with hematological malignancy received cyclophosphamide (120 mg/kg), fractionated total body irradiation (12 Gy), oral cyclosporin, and an HLA-identical sibling marrow transplant depleted of T cells by incubation with the monoclonal **antibody** antiHuLy-m1 (CD2) and rabbit complement with (five patients) or without (11 patients) anti-HuLy-m8 (CD8). These 16 patients were compared historically to 84 patients with hematological malignancy receiving cyclophosphamide (120 mg/kg), fractionated total body irradiation (12 or 14 Gy), oral cyclosporin, and unmanipulated HLA-identical sibling marrow, for parameters of engraftment and graft-versus-host disease (GVHD). Graft failure occurred in one of the 16 T-cell depleted recipients and in one of the 84 **non-depleted** recipients. Engraftment was slightly but significantly slower in the T-cell depleted group and bacterial infections significantly more frequent and severe than in the unmanipulated group. There was a suggestion that the severity of acute GVHD was reduced in those receiving T depleted marrow. Randomized trials will be necessary to determine if marrow T-cell depletion results in superior long-term leukemia-free survival.

9/7/5 (Item 1 from file: 72)

DIALOG(R) File 72:EMBASE

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8675097 EMBASE No: 92355607

Anti-CD4 monoclonal **antibodies** in therapy: Creation of nonclassical tolerance in the adult

Shizuru J.A.; Alters S.E.; Fathman C.G.

Stanford Univ. School of Medicine, Div. of Rheumatology and Immunology, Stanford, CA 94305 USA

IMMUNOL. REV. (Denmark) , 1992, -/129 (105-130)

CODEN: IMRED ISSN: 0105-2896 ADONIS ORDER NUMBER: 010528969200046X

LANGUAGES: English SUMMARY LANGUAGES: English

We have described the studies from our laboratory which demonstrate that depleting anti-CD4 mAb induce tolerance to foreign antigens in adult, euthymic animals. Further, we have proposed that such tolerance occurs as a result of new thymic migrants encountering antigens in the periphery. However, these conclusions can be considered only partial since we (Song et al. in press) and others have shown that depletion of T cells per se does not permit tolerance. For example, anti-Thy-1 or anti-Lyt-1 are themselves immunosuppressive and able to deplete T cells, yet they elicit strong anti-globulin responses against themselves and do not permit tolerance to be induced either to transplants or administered soluble protein antigen. We have recently found that while the combination of anti-CD4 and anti-CD8 mAb allows long-term survival of allografted islets in mice, anergy in the relevant T-cell subsets was not found (in contrast to what is found with anti-CD4 mAb treatment alone) (Song et al. in press). In this instance, long-term survival was probably the result of changes in graft immunogenicity (i.e., migration of passenger leukocytes) since the kinetics of repopulation were much delayed in the anti-CD4 and -CD8 treated mice. As discussed elsewhere in this volume, interesting studies from several laboratories suggest that **non-depleting** anti-CD4 mAb can generate unresponsiveness in a variety of systems. In reviewing the literature it is clear that the success of **non-depleting** reagents appears to be dependent upon the model system tested. For example, although depleting and **nondepleting** CD4 mAb regimens produced

comparable prolongation of cultured fetal pancreas allografts in mice (Charlton and Mandel), almost total elimination of circulating CD4+ cells did not prevent acute rejection of murine skin grafts (Auchincloss et al. 1988). This heterogeneity is not surprising given the multiple functional roles of the CD4 molecule and the cells that bear this molecule. In addition to depletion, **antibodies** directed against CD4 can potentially affect CD4+ cell function by (1) direct blockade or failure to augment the formation of the TCR-antigen/MHC ternary complex or (2) by transmitting a negative signal to the CD4 T cell or interfering with normal signal transduction mechanisms. Undoubtedly, it is a combination of mechanisms that allows these **antibodies** their immunosuppressive effects. What can be said with certainty is that these **antibodies** will continue to be important tools for understanding the molecular and cellular basis of the immune response, and will soon emerge as invaluable therapeutic agents in the clinical arena.

9/7/6 (Item 2 from file: 72)
DIALOG(R) File 72:EMBASE
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8183556 EMBASE No: 91209639
Monoclonal **antibody** therapy for the induction of transplantation tolerance
Cobbold S.P.
Division of Immunology, Cambridge University Department of Pathology,
Tennis Court Road, Cambridge CB1 2QP United Kingdom
IMMUNOL. LETT. (Netherlands) , 1991, 29/1-2 (117-122)
CODEN: IMLED ISSN: 0165-2478 ADONIS ORDER NUMBER: 016524789100175N
LANGUAGES: English
There are three ways in which monoclonal **antibodies** could be used to facilitate the induction of tolerance to foreign tissues after organ transplantation. First, depleting monoclonal **antibodies** could be directed against the T cells responsible, thereby reducing their number and acting to non-specifically immunosuppress the patient. This is generally not sufficient to allow tolerance induction in the T cells which repopulate the periphery. Second, depleting monoclonal **antibodies** could be used to remove donor passenger leukocytes and antigen-presenting cells from the donor organ, which may both reduce immunogenicity and increase the chance of tolerance induction. Third, **non-depleting**, but functionally blocking, monoclonal **antibodies** to T cell molecules such as CD4 and CD8 can allow the specific induction of transplantation tolerance in mouse models, an approach which might be applicable to man, not only for organ transplantation, but also in the treatment of autoimmune diseases. These three approaches are, in time, likely to complement each other in clinical practice. Monoclonal **antibodies** can be tailored to each approach by choosing appropriate specificities and isotypes, and further refinements can be made where necessary by making monovalent or **humanised antibodies**. The application of each of these approaches to clinical therapy is described.

9/7/7 (Item 3 from file: 72)
DIALOG(R) File 72:EMBASE
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8013038 EMBASE No: 91038466
Induction of tolerance in peripheral T cells with monoclonal **antibodies**
Qin S.; Wise M.; Cobbold S.P.; Leong L.; Kong Y.-C.M.; Parnes J.R.;
Waldmann H.
Division of Immunology, Department of Pathology, Cambridge University,
Cambridge CB2 2QQ United Kingdom
EUR. J. IMMUNOL. (Germany, Federal Republic of) , 1990, 20/12 (2737-2745)
CODEN: EJIMA ISSN: 0014-2980

LANGUAGES: English

Our goal has been to develop ways to tolerize the mature immune system to any defined antigen. In this report we show that peripheral (post-thymic) T cells of mice can become tolerant to a range of antigens (**human** and **rat immunoglobulins**, and bone marrow and skin grafts that differ at multiple minor transplantation antigens). In the case of **human** gamma globulin (HGG), this required that the antigen be given under the cover of a short course of **non-depleting** anti-CD4 **antibody**, while for tolerance to skin and marrow grafts anti-CD8 **antibody** was also required. Tolerance to HGG could be reinforced by repeated injections of HGG, but was lost in the absence of any further exposure to antigen. This reversal of tolerance with time was due to new T cells being exported from the thymus, as it was not observed in tolerized, adult thymectomized mice. In contrast, tolerance to marrow and skin grafts was permanent, presumably because the established grafts acted as a continuous source of antigen to reinforce the tolerant state. Tolerance could not be broken by the infusion of unprimed spleen cells and in one example (tolerance to Mls-1a) there was clear evidence that specific peripheral T cells were anergic. We propose that anergic cells may themselves participate in reinforcing the tolerant state by competing at sites of antigen presentation.

9/7/8 (Item 1 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

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09426656 98149797

Treatment of recalcitrant plaque psoriasis with a **humanized non-depleting antibody** to CD4.

Bachelez H; Flageul B; Dubertret L; Fraitag S; Grossman R; Brousse N; Poisson D; Knowles RW; Wacholtz MC; Haverty TP; Chatenoud L; Bach JF Service Dermatologie, Hopital Saint-Louis, Paris, France.

J Autoimmun (ENGLAND) Feb 1998, 11 (1) p53-62, ISSN 0896-8411

Journal Code: ADL

Languages: ENGLISH

Document type: JOURNAL ARTICLE

The presence of activated CD4(+) T lymphocytes in psoriatic skin plaques suggests an immune-mediated pathogenesis for the disease. Six patients with recalcitrant plaque psoriasis (PASI>12) received a **humanized non-depleting** monoclonal **antibody** to CD4 (ORTHOCLONE OKT(R)cdr4a). The **antibody** was well tolerated. Four weeks from treatment, the mean decrease in PASI score was 46%. In three patients disease remission was prolonged for up to 6 months and, in one case, up to 1 year post-treatment. In all patients, circulating CD4+ T-cell counts remained normal and peripheral OKTcdr4a-coated CD4+ lymphocytes were detected up to 10 days after **antibody** infusion. These results point to the relevance of CD4+ lymphocytes in psoriasis. They also emphasize that depletion of CD4+ cells is not mandatory to achieve therapeutic effectiveness. Copyright 1998 Academic Press Limited.

9/7/9 (Item 2 from file: 154)

DIALOG(R) File 154: MEDLINE(R)

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06603039 91370929

Reprogramming the immune system for tolerance with monoclonal **antibodies**.

Cobbold SP; Qin SX; Waldmann H
Department of Pathology, Cambridge University, UK.
Semin Immunol (UNITED STATES) Nov 1990, 2 (6) p377-87, ISSN 1044-5323
Journal Code: A61

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

Monoclonal **antibodies** to CD4, **CD8** and CD11a can be used in vivo either to deplete or functionally block T cells to create a tolerance permissive environment. Short courses of **non-depleting** CD4 and **CD8 antibodies** were used to induce tolerance separately in CD4+ and CD8+ T cells either to foreign **immunoglobulins**, bone marrow, or skin grafts. Tolerance was obtained to minor (non-MHC) transplantation antigens without T cell depletion even in actively sensitized mice, or to MHC plus minor antigens presented directly by skin grafts using combinations of depleting followed by blockading CD4 and **CD8 antibodies**. In all cases, tolerance was specific to the antigen/tissue given under cover of **antibody** treatment, and in one example it could be shown that T cells directed to MLS-1a had been forced into an anergic state. This induction of tolerant, anergic T cells in the periphery is able to explain many of the features associated with tolerance, not only in the model systems using foreign antigens, but also in the normal regulation of anti-self responses and its failure in autoimmune diseases. It is our new found ability to use antigen under the cover of **antibody** treatment to accurately control the pattern of tolerant T cells in vivo that we refer to by using the term 'reprogramming'. We also describe the clinical treatment of one patient with an autoimmune vasculitis based on the ideas developed from the mouse models. (47 Refs.)

9/7/10 (Item 1 from file: 351)
 DIALOG(R) File 351:DERWENT WPI
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008503137

WPI Acc No: 91-007221/199101

Non-depleting CD4 and **CD8** monoclonal **antibodies** -
 for inducting tolerance to foreign antigens in transplant rejection,
 auto-immune disease, etc

Patent Assignee: COBBOLD S P (COBB-I); WALDMANN H (WALD-I); WELLCOME FOUND LTD (WELL); GLAXO WELLCOME INC (GLAX)

Inventor: COBBOLD S P; WALDMANN H

Number of Countries: 025 Number of Patents: 015

Patent Family:

Patent No	Kind	Date	Applicat No	Kind	Date	Main IPC	Week
WO 9015152	A	19901213					199101 B
PT 94214	A	19910208					199109
AU 9057258	A	19910107					199115
EP 474691	A	19920318	EP 90908270	A	19900531		199212
ZA 9004174	A	19920226	ZA 904174	A	19900530		199213
DD 296843	A5	19911219	DD 341218	A	19900531	A61K-039/395	199221
JP 4505919	W	19921015	JP 90508030	A	19900531	A61K-039/395	199248
			WO 90GB840	A	19900531		
HU 61341	T	19921230	HU 905134	A	19900531	C12P-021/08	199306
			WO 90GB840	A	19900531		
AU 657255	B	19950309	AU 9057258	A	19900531	C12P-021/08	199520
EP 474691	B1	19961113	EP 90908270	A	19900531	C12P-021/08	199650
			WO 90GB840	A	19900531		
DE 69029134	E	19961219	DE 629134	A	19900531	C12P-021/08	199705
			EP 90908270	A	19900531		
			WO 90GB840	A	19900531		
ES 2096588	T3	19970316	EP 90908270	A	19900531	C12P-021/08	199718
NZ 233889	A	19970624	NZ 233889	A	19900531	A61K-039/395	199732
BR 1100287	A3	19970916	BR 971100287	A	19970415	C12P-021/08	199744
US 5690933	A	19971125	US 91768868	A	19910727	A61K-039/395	199802
			US 9347344	A	19930329		
			US 94181170	A	19940113		
			US 94289532	A	19940812		

Priority Applications (No Type Date): GB 8912497 A 19890531
 Cited Patents: 4.Jnl.Ref

Patent Details:

Patent	Kind	La	Pa	Filing	Notes	Application	Patent
WO							
9015152	A						
Designated States (National): AU CA FI HU JP KR US							
Designated States (Regional): AT BE CH DE DK ES FR GB IT LU NL SE							
EP	474691	A		44			
Designated States (Regional): AT BE CH DE DK ES FR GB IT LI LU NL SE							
ZA	9004174	A		57			
JP	4505919	W		19	Based on	WO	9015152
HU	61341	T			Based on	WO	9015152
AU	657255	B			Previous Publ.	AU	9057258
					Based on	WO	9015152
EP	474691	B1	E	32	Based on	WO	9015152
Designated States (Regional): AT BE CH DE DK ES FR GB IT LI LU NL SE							
DE	69029134	E			Based on	EP	474691
					Based on	WO	9015152
ES	2096588	T3			Based on	EP	474691
US	5690933	A		23	Cont of	US	91768868
					Cont of	US	9347344
					Cont of	US	94181170

Abstract (Basic): WO 9015152 A

Non depleting CD4 and CD8 monoclonal antibodies are claimed for use in inducing tolerance to an antigen. The use of these antibodies and packs contg. them are also claimed. The prods. may also contain a depleting CD4 monoclonal antibody and/or a depleting CD8 monoclonal antibody.

Single dose for a human is 1-400mg (esp. 3-30mg) of antibody. Admin. is parenteral e.g. intravenous.

USE/ADVANTAGE - For producing tolerance to foreign immunoglobulins, bone marrow and skin grafts. To treat autoimmune diseases without the need for long term chemotherapy and to produce tolerance to therapeutic polypeptides such as interferon, IL-II or TNF. (44pp Dwg. No. 0/13)

Abstract (Equivalent): EP 474691 B

Use of a non-depleting anti-CD4 monoclonal antibody, ie an antibody which causes depletion of fewer than 50% of CD4+ T-cells from the periphery as measured by changes in the peripheral blood lymphocyte numbers, for the manufacture of a medicament for the induction of a state of immunological tolerance to an antigen by a method which comprises administering said non-depleting anti-CD4 monoclonal antibody to a subject together with a non-depleting anti-CD8 monoclonal antibody, ie an antibody which causes depletion of fewer than 50% of CD8+ T-cells from the periphery as measured by changes in the peripheral blood lymphocyte numbers, to induce an immunological tolerance permissive environment within said subject by means of said antibodies in the presence of said antigen.

(Dwg. 0/11b)

Abstract (Equivalent): US 5690933 A

Non depleting CD4 and CD8 monoclonal antibodies are claimed for use in inducing tolerance to an antigen. The use of these antibodies and packs contg. them are also claimed. The prods. may also contain a depleting CD4 monoclonal antibody and/or a depleting CD8 monoclonal antibody.

Single dose for a human is 1-400mg (esp. 3-30mg) of antibody. Admin. is parenteral e.g. intravenous.

USE/ADVANTAGE - For producing tolerance to foreign immunoglobulins, bone marrow and skin grafts. To treat autoimmune diseases without the need for long term chemotherapy and to produce tolerance to therapeutic polypeptides such as interferon, IL-II or TNF.

Dwg. 0/13b

Derwent Class: B04; D16

International Patent Class (Main): A61K-039/395; C12P-021/08

International Patent Class (Additional): A61K-037/02; A61K-039/39;

C07K-015/28; C07K-016/24; C07K-016/42; C12N-015/00; G01N-000/00
? ds

Set Items Description
S1 312 (NON(W) DEPLET? OR NONDEPLET?) AND (ANTIBOD? OR IMMUNOGLOBU-
 LIN?)
S2 224 S1 AND CD4
S3 67 S2 AND HUMAN?
S4 45 RD S3 (unique items)
S5 95 S1 AND CD8
S6 42 RD S5 (unique items)
S7 6 S2 AND REVIEW?
S8 10 S6 AND HUMAN?
S9 10 RD S8 (unique items)
? s s2 and py=1988

 224 S2
 2151700 PY=1988
S10 3 S2 AND PY=1988
? rd s10

>>>Duplicate detection is not supported for File 351.

>>>Records from unsupported files will be retained in the RD set.
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 S11 1 RD S10 (unique items)
? t s11/3/all

11/3/1 (Item 1 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

7010730 BIOSIS Number: 87071251
ADOPTIVE IMMUNITY IN IMMUNE-DEFICIENT SCID-SCID MICE I. DIFFERENTIAL
REQUIREMENTS OF NAIVE AND PRIMED LYMPHOCYTES FOR CD4-POSITIVE T CELLS
DURING REJECTION OF MINOR HISTOCOMPATIBILITY ANTIGEN-DISPARATE SKIN GRAFTS
ROOPENIAN D C; ANDERSON P S
JACKSON LAB., BAR HARBOR, ME 04609.
TRANSPLANTATION (BALTIMORE) 46 (6). 1988. 899-904. CODEN: TRPLA
Full Journal Title: TRANSPLANTATION (Baltimore)
Language: ENGLISH
? s s2 and py=1989

 224 S2
 2238453 PY=1989
S12 9 S2 AND PY=1989
? rd s12

>>>Duplicate detection is not supported for File 351.

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? t s13/3/all

13/3/1 (Item 1 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

7185784 BIOSIS Number: 88108529
ENGAGEMENT OF CD-4 AND CD-8 ACCESSORY MOLECULES IS REQUIRED FOR T CELL
MATURATION

RAMSDELL F; FOWLKES B J
LAB. CELLULAR MOLECULAR IMMUNOL., NIAID, NIH, BUILDING 4, ROOM 111,
BETHESDA, MD 20892.
J IMMUNOL 143 (5). 1989. 1467-1471. CODEN: JOIMA
Full Journal Title: Journal of Immunology
Language: ENGLISH

13/3/2 (Item 2 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

7104252 BIOSIS Number: 88026997
AN INCREASE IN THE SURVIVAL OF MURINE H-2-MISMATCHED CULTURED FETAL
PANCREAS ALLOGRAFTS USING DEPLETING OR **NONDEPLETING ANTI-CD4**
MONOCLONAL **ANTIBODIES** AND A FURTHER INCREASE WITH THE ADDITION OF
CYCLOSPORINE
BURKHARDT K; CHARLTON B; MANDEL T E
TRANSPLANTATION UNIT, WALTER AND ELIZA HALL INST. MED. RES., POST OFFICE,
ROYAL MELBOURNE HOSP., VICTORIA 3050, AUST.
TRANSPLANTATION (BALTIMORE) 47 (5). 1989. 771-775. CODEN: TRPLA
Full Journal Title: TRANSPLANTATION (Baltimore)
Language: ENGLISH

13/3/3 (Item 3 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

7010433 BIOSIS Number: 87070954
T-CELL-MEDIATED PROTECTION OF MICE AGAINST VIRULENT
MYCOBACTERIUM-TUBERCULOSIS
LEVETON C; BARNASS S; CHAMPION B; LUCAS S; DE SOUZA B; NICOL M; BANERJEE
D; ROOK G
DEP. MED. MICROBIOL., UNIV. COLL., LONDON W1P 7PP, U.K.
INFECT IMMUN 57 (2). 1989. 390-395. CODEN: INFIB
Full Journal Title: Infection and Immunity
Language: ENGLISH
? s s2 and py=1987

224 S2
2075095 PY=1987
S14 2 S2 AND PY=1987
? rd s14

>>>Duplicate detection is not supported for File 351.

>>>Records from unsupported files will be retained in the RD set.
...completed examining records
S15 1 RD S14 (unique items)
? t s15/3/all

15/3/1 (Item 1 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

5934551 BIOSIS Number: 84067116
CD4 POSITIVE T CELLS APPEAR CAPABLE OF INITIATING GRAFT-VERSUS-HOST
DISEASE ACROSS NON-MAJOR HISTOCOMPATIBILITY COMPLEX MHC BARRIERS IN MAN
ATKINSON K; COOLEY M; FARRELLY H; O'FLAHERTY E; ASHBY M; BIGGS J
DEP. HAEMATOL., ST VINCENT'S HOSP., DARLINGHURST, NSW 2010, AUSTRALIA.
BONE MARROW TRANSPLANT 2 (1). 1987. 79-84. CODEN: BMTRE
Full Journal Title: Bone Marrow Transplantation
Language: ENGLISH

? s s2 and py=1990

224 S2
2373119 PY=1990
S16 11 S2 AND PY=1990
? rd s16

>>>Duplicate detection is not supported for File 351.

>>>Records from unsupported files will be retained in the RD set.
...completed examining records
S17 6 RD S16 (unique items)
? t s17/7/all

17/7/1 (Item 1 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
(c) 1998 BIOSIS. All rts. reserv.

8167526 BIOSIS Number: 91088526
THE INDUCTION OF SKIN GRAFT TOLERANCE IN MAJOR HISTOCOMPATIBILITY
COMPLEX-MISMATCHED OR PRIMED RECIPIENTS PRIMED T CELLS CAN BE TOLERIZED IN
THE PERIPHERY WITH ANTI-**CD4** AND ANTI-CD8 **ANTIBODIES**
COBBOLD S P; MARTIN G; WALDMANN H
DIV. IMMUNOL., CAMBRIDGE UNIV., DEP. PATHOL., LEVEL 3 LAB. BLOCK, NEW
ADDENBROOKES HOSP., CAMBRIDGE CB2 2QQ, GREAT BRITIAN.
EUR J IMMUNOL 20 (12). 1990. 2747-2756. CODEN: EJIMA
Full Journal Title: European Journal of Immunology
Language: ENGLISH
Mice given short courses of anti-**CD4** and anti-CD8 monoclonal
antibodies became tolerant of allogeneic skin grafted at the same
time. Tolerance could be obtained without T cell depletion across multiple
minor antigen mismatches, both in native and primed animals, demonstrating
that peripheral T cells could be tolerized, even if they had been
previously activated. Where donor and recipient were incompatible across
the whole major histocompatibility complex, specific tolerance could be
achieved by using a combination of depleting following by **non-**
depleting antibodies, where each alone was unsuccessful.
Although mice clearly tolerated their original skin grafts, we observed in
some strain combinations that a second fresh, but genotypically indentical
graft, was slowly rejected. Such mice also possessed T cells which could
proliferate to donor-type stimulator cells in vitro. Whatever the
mechanisms, we have demonstrated that operational transplantation tolerance
can be achieved with simple, non-toxic **antibody** therapy. The
introduction of comparable tolerance-inducing regimens in clinical organ
transplantation could obviate the need for long-term immunosuppression and
its unfortunate side effects.

17/7/2 (Item 2 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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8095291 BIOSIS Number: 91016291
RESISTANCE TO INFECTION BY HIV-1 OF PERIPHERAL BLOOD MONONUCLEAR CELLS
FROM HIV-1-INFECTED PATIENTS IS PROBABLY MEDIATED BY NEUTRALIZING
ANTIBODIES
TREMBLAY M; NUMAZAKI K; LI X; GORNITSKY M; HISCOTT J; WAINBERG M A
MCGILL AIDS CENTRE JEWISH GENERAL HOSP., 3755 COTE STE-CATHERINE ROAD,
MONTREAL, QUEBEC H3T 1E2, CAN.
J IMMUNOL 145 (9). 1990. 2896-2901. CODEN: JOIMA
Full Journal Title: Journal of Immunology
Language: ENGLISH
We have investigated whether PBMC of HIV-1-seropositive subjects are as
susceptible to in vitro infection by HIV-1 as are PBMC from seronegative

controls. Accordingly, stimulated PBMC from 19 HIV-1-infected subjects were inoculated with four different variants of HIV-1. None of these cultures produced either detectable quantities of viral reverse transcriptase activity or p24 Ag following inoculation with HIV-1. In contrast, in five of six cases in which these PBMC were depleted of B cells by **antibody** plus complement prior to viral inoculation, the presence of viral reverse transcriptase and p24 Ag was detected. The presence of normal levels of CD4 Ag at the surface of the CD4+ cells in these populations was established by flow cytometry. Analysis by an immunoblot assay revealed that anti-HIV **antibodies** were present in the sera obtained from these infected donors; in addition, 7 of 10 culture fluids derived from the **nondepleted** PBMC were shown to contain virus-neutralizing **antibodies**. Cultures which were depleted of B cells did not contain detectable levels of antiviral **antibodies**. Confirmation that the virus produced by the PBMC which had been depleted of B cells was of the strain used to infect the cultures, rather than that which initially caused patient infection, was provided on the basis of differential susceptibility to **antibody** neutralization. These results suggest that **antibodies** produced by B cells in cultures of PBMC from seropositive donors may restrict infection by HIV-1 of such cultures under laboratory conditions.

17/7/3 (Item 3 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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7529494 BIOSIS Number: 39042101
A **NONDEPLETING RAT CD4 MONOCLONAL ANTIBODY MAB INHIBITS CD4-POSITIVE SUPPRESSOR-MEDIATED RESISTANCE TO MURINE EXPERIMENTAL AUTO-IMMUNE THYROIDITIS EAT IN-VIVO**
NABOZNY G H; COBBOLD S; WALDMANN H; KONG Y M
WAYNE STATE UNIV. SCH. MED., DETROIT, MICH. 48201.
JOINT MEETING OF THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY, AND THE AMERICAN ASSOCIATION OF IMMUNOLOGISTS, NEW ORLEANS, LOUISIANA, USA, JUNE 4-7, 1990. FASEB (FED AM SOC EXP BIOL) J 4 (7). 1990. A2099. CODEN: FAJ0E
Language: ENGLISH

17/7/4 (Item 1 from file: 72)
DIALOG(R) File 72:EMBASE
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8013038 EMBASE No: 91038466
Induction of tolerance in peripheral T cells with monoclonal **antibodies**
Qin S.; Wise M.; Cobbold S.P.; Leong L.; Kong Y.-C.M.; Parnes J.R.; Waldmann H.

Division of Immunology, Department of Pathology, Cambridge University, Cambridge CB2 2QQ United Kingdom

EUR. J. IMMUNOL. (Germany, Federal Republic of) , 1990, 20/12 (2737-2745)

CODEN: EJIMA ISSN: 0014-2980
LANGUAGES: English

Our goal has been to develop ways to tolerize the mature immune system to any defined antigen. In this report we show that peripheral (post-thymic) T cells of mice can become tolerant to a range of antigens (human and rat **immunoglobulins**, and bone marrow and skin grafts that differ at multiple minor transplantation antigens). In the case of human gamma globulin (HGG), this required that the antigen be given under the cover of a short course of **non-depleting anti-CD4 antibody**, while for tolerance to skin and marrow grafts anti-CD8 **antibody** was also required. Tolerance to HGG could be reinforced by repeated injections of HGG, but was lost in the absence of any further exposure to antigen.

This reversal of tolerance with time was due to new T cells being exported from the thymus, as it was not observed in tolerized, adult thymectomized mice. In contrast, tolerance to marrow and skin grafts was permanent, presumably because the established grafts acted as a continuous source of antigen to reinforce the tolerant state. Tolerance could not be broken by the infusion of unprimed spleen cells and in one example (tolerance to Mls-1a) there was clear evidence that specific peripheral T cells were anergic. We propose that anergic cells may themselves participate in reinforcing the tolerant state by competing at sites of antigen presentation.

17/7/5 (Item 1 from file: 154)
DIALOG(R) File 154: MEDLINE(R)
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06603039 91370929
Reprogramming the immune system for tolerance with monoclonal antibodies.
Cobbold SP; Qin SX; Waldmann H
Department of Pathology, Cambridge University, UK.
Semin Immunol (UNITED STATES) Nov 1990, 2 (6) p377-87, ISSN
1044-5323 Journal Code: A61
Languages: ENGLISH
Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL
Monoclonal antibodies to CD4, CD8 and CD11a can be used in vivo either to deplete or functionally block T cells to create a tolerance permissive environment. Short courses of non-depleting CD4 and CD8 antibodies were used to induce tolerance separately in CD4+ and CD8+ T cells either to foreign immunoglobulins, bone marrow, or skin grafts. Tolerance was obtained to minor (non-MHC) transplantation antigens without T cell depletion even in actively sensitized mice, or to MHC plus minor antigens presented directly by skin grafts using combinations of depleting followed by blockading CD4 and CD8 antibodies. In all cases, tolerance was specific to the antigen/tissue given under cover of antibody treatment, and in one example it could be shown that T cells directed to MLS-1a had been forced into an anergic state. This induction of tolerant, anergic T cells in the periphery is able to explain many of the features associated with tolerance, not only in the model systems using foreign antigens, but also in the normal regulation of anti-self responses and its failure in autoimmune diseases. It is our new found ability to use antigen under the cover of antibody treatment to accurately control the pattern of tolerant T cells in vivo that we refer to by using the term 'reprogramming'. We also describe the clinical treatment of one patient with an autoimmune vasculitis based on the ideas developed from the mouse models. (47 Refs.)

17/7/6 (Item 1 from file: 351)
DIALOG(R) File 351: DERWENT WPI
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008503137
WPI Acc No: 91-007221/199101
Non-depleting CD4 and CD8 monoclonal antibodies -
for inducing tolerance to foreign antigens in transplant rejection,
auto-immune disease, etc
Patent Assignee: COBBOLD S P (COBB-I); WALDMANN H (WALD-I); WELLCOME FOUND
LTD (WELL); GLAXO WELLCOME INC (GLAX)
Inventor: COBBOLD S P; WALDMANN H
Number of Countries: 025 Number of Patents: 015
Patent Family:
Patent No Kind Date Applicat No Kind Date Main IPC Week
WO 9015152 A 19901213 199101 B

PT 94214	A	19910208		199109	
AU 9057258	A	19910107		199115	
EP 474691	A	19920318	EP 90908270	A 19900531	199212
ZA 9004174	A	19920226	ZA 904174	A 19900530	199213
DD 296843	A5	19911219	DD 341218	A 19900531	A61K-039/395 199221
JP 4505919	W	19921015	JP 90508030	A 19900531	A61K-039/395 199248
			WO 90GB840	A 19900531	
HU 61341	T	19921230	HU 905134	A 19900531	C12P-021/08 199306
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			WO 90GB840	A 19900531	
DE 69029134	E	19961219	DE 629134	A 19900531	C12P-021/08 199705
			EP 90908270	A 19900531	
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ES 2096588	T3	19970316	EP 90908270	A 19900531	C12P-021/08 199718
NZ 233889	A	19970624	NZ 233889	A 19900531	A61K-039/395 199732
BR 1100287	A3	19970916	BR 971100287	A 19970415	C12P-021/08 199744
US 5690933	A	19971125	US 91768868	A 19910727	A61K-039/395 199802
			US 9347344	A 19930329	
			US 94181170	A 19940113	
			US 94289532	A 19940812	

Priority Applications (No Type Date): GB 8912497 A 19890531

Cited Patents: 4.Jnl.Ref

Patent Details:

Patent	Kind	Ln	Pg	Filing Notes	Application	Patent
WO 9015152	A					
				Designated States (National): AU CA FI HU JP KR US		
				Designated States (Regional): AT BE CH DE DK ES FR GB IT LU NL SE		
EP 474691	A		44			
				Designated States (Regional): AT BE CH DE DK ES FR GB IT LI LU NL SE		
ZA 9004174	A		57			
JP 4505919	W		19	Based on	WO 9015152	
HU 61341	T			Based on	WO 9015152	
AU 657255	B			Previous Publ.	AU 9057258	
				Based on	WO 9015152	
EP 474691	B1	E	32	Based on	WO 9015152	
				Designated States (Regional): AT BE CH DE DK ES FR GB IT LI LU NL SE		
DE 69029134	E			Based on	EP 474691	
				Based on	WO 9015152	
ES 2096588	T3			Based on	EP 474691	
US 5690933	A		23	Cont of	US 91768868	
				Cont of	US 9347344	
				Cont of	US 94181170	

Abstract (Basic): WO 9015152 A

Non depleting CD4 and CD8 monoclonal antibodies are claimed for use in inducing tolerance to an antigen. The use of these antibodies and packs contg. them are also claimed. The prods. may also contain a depleting CD4 monoclonal antibody and/or a depleting CD8 monoclonal antibody.

Single dose for a human is 1-400mg (esp. 3-30mg) of antibody. Admin. is parenteral e.g. intravenous.

USE/ADVANTAGE - For producing tolerance to foreign immunoglobulins, bone marrow and skin grafts. To treat autoimmune diseases without the need for long term chemotherapy and to produce tolerance to therapeutic polypeptides such as interferon, IL-II or TNF. (44pp Dwg.No.0/13)

Abstract (Equivalent): EP 474691 B

Use of a non-depleting anti-CD4 monoclonal antibody, ie an antibody which causes depletion of fewer than 50% of CD4+ T-cells from the periphery as measured by changes in the peripheral blood lymphocyte numbers, for the manufacture

of a medicament for the induction of a state of immunological tolerance to an antigen by a method which comprises administering said **non-depleting** anti-**CD4** monoclonal **antibody** to a subject together with a **non-depleting** anti-**CD8** monoclonal **antibody**, ie an **antibody** which causes depletion of fewer than 50% of **CD8+** T-cells from the periphery as measured by changes in the peripheral blood lymphocyte numbers, to induce an immunological tolerance permissive environment within said subject by means of said **antibodies** in the presence of said antigen.

(Dwg.0/11b

Abstract (Equivalent): US 5690933 A

Non depleting CD4 and **CD8** monoclonal **antibodies** are claimed for use in inducing tolerance to an antigen. The use of these **antibodies** and packs contg. them are also claimed. The prods. may also contain a depleting **CD4** monoclonal **antibody** and/or a depleting **CD8** monoclonal **antibody**.

Single dose for a human is 1-400mg (esp. 3-30mg) of **antibody**. Admin. is parenteral e.g. intravenous.

USE/ADVANTAGE - For producing tolerance to foreign **immunoglobulins**, bone marrow and skin grafts. To treat autoimmune diseases without the need for long term chemotherapy and to produce tolerance to therapeutic polypeptides such as interferon, IL-II or TNF.

Dwg.0/13b

Derwent Class: B04; D16

International Patent Class (Main): A61K-039/395; C12P-021/08

International Patent Class (Additional): A61K-037/02; A61K-039/39; C07K-015/28; C07K-016/24; C07K-016/42; C12N-015/00; G01N-000/00
? t s13/7/2

13/7/2 (Item 2 from file: 55)
DIALOG(R) File 55:BIOSIS PREVIEWS(R)
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7104252 BIOSIS Number: 88026997

AN INCREASE IN THE SURVIVAL OF MURINE H-2-MISMATCHED CULTURED FETAL PANCREAS ALLOGRAFTS USING DEPLETING OR **NONDEPLETING ANTI-CD4** MONOCLONAL **ANTIBODIES** AND A FURTHER INCREASE WITH THE ADDITION OF CYCLOSPORINE

BURKHARDT K; CHARLTON B; MANDEL T E
TRANSPLANTATION UNIT, WALTER AND ELIZA HALL INST. MED. RES., POST OFFICE, ROYAL MELBOURNE HOSP., VICTORIA 3050, AUST.

TRANSPLANTATION (BALTIMORE) 47 (5). 1989. 771-775. CODEN: TRPLA
Full Journal Title: TRANSPLANTATION (Baltimore)

Language: ENGLISH
Depletion of **CD4+** T lymphocytes with monoclonal **antibodies** (mAbs) has been shown to prolong allograft survival in mice. In this study, two rat anti-**CD4** mAbs, H129.19 and GK1.5, were administered either alone or in combination with cyclosporine (CsA) to recipients of MHC-mismatched (H-2k to H-2d) cultured fetal pancreas allografts to determine their effect on graft survival. When compared with control mice, splenic **CD4+** cells of GK1.5-treated mice were depleted by > 95%, but in H129.19-treated mice no depletion of **CD4** + cells occurred. Instead, rat Ig was present on the surface of **CD4** + cells in H129.19-treated mice. Anti-**CD4** therapy with either H129.19 or GK1.5 prolonged fetal pancreas allograft survival to a similar extent, but did not lead to indefinite survival. Blockade of the **CD4** antigen by the mAb H129.19 was as effective as the depletion of **CD4+** cells by GK1.5 in prolonging allograft survival. Rejection of grafts by day 28 posttransplantation occurred in the absence of **CD4** + cells, as determined by both flow cytometric examination of spleen cells and immunoperoxidase staining of the graft site. CsA alone did not prolong graft survival, but its addition to either H129.19 or GK1.5 mAb treatment significantly increased the survival rate of grafts at 28 days compared

with mAb treatment alone. These results suggest that **CD4+** cell depletion is not essential for effective anti-**CD4** mAb therapy-and, further, that CSA may have a direct inhibitory effect on CD8+ cells during allograft rejection.